U. S. DEPARTMENT OF COMMERCE

HENRY A. WALLACE, Secretary

NATIONAL BUREAU OF STANDARDS

E. U. CONDON, Director

CIRCULAR OF THE BUREAU OF STANDARDS C450

UNDERGROUND CORROSION

[Issued November 27, 1945]



PREFACE

Bureau Circular C401, Abstracts and Summaries of Bureau of Standards Publications on Stray-Current Electrolysis, was published in 1933. This gives the essential conclusions drawn from the work of the National Bureau of Standards on stray-current electrolysis during a period of 15 years, as presented in 17 Bureau Technologic Papers, most of which are available only in reference libraries.

As corrosion due to stray-current electrolysis became less important, attention was turned to underground corrosion due to causes other than stray currents. Although this work has been interrupted by the necessity of giving full attention to war work, the subject continues to be one of major importance. A considerable number of Bureau publications have been issued, but most of them are also out of print

and available only in reference libraries.

There is a continuing demand for technical information relating to the problem of underground corrosion. Because of the extensive installation and repair of underground structures that will occur in the postwar period the Bureau feels that the issuance of a publication giving under one cover the research results obtained thus far, together with a general discussion of the corrosion problem and the conclusions to be drawn from the data, will serve a useful purpose. These conclusions are supplemented by summaries of the results of the more important investigations of underground-corrosion phenomena carried on in America and Europe.

The manufacturers who supplied the materials tested have rendered valuable assistance in planning the tests and in the interpretation of the results. The cooperation of the public utilities and the technical societies

has also been of great value in the work.

The material for this Circular is based on the results obtained by many members of the Bureau's Underground Corrosion Section over a period of several years, and the use of their data is acknowledged by the author.

· E. U. CONDON, Director.

UNDERGROUND CORROSION

By Kirk H. Logan

ABSTRACT

This Circular is an assembly of the results of the National Bureau of Standards investigations of underground corrosion which began in 1922. The fundamental causes and processes of underground corrosion are the same as those occurring in the air or water, but their relative values are different. Corrosion in soils is the result of soil characteristics and conditions but these are too numerous and complex to permit a satisfactory correlation of corrosion with any single soil property.

The results of the tests of ferrous materials given in this Circular indicate that the commonly used ferrous pipe materials do not differ greatly in their resistance to soils and that their apparent relative merits are either accidental or dependent on soil conditions. Low-alloy ferrous materials lose weight more slowly than unalloyed ferrous materials, but are penetrated by corrosion as rapidly. Alloys high in nickel

and chromium are very resistant to corrosion.

In most of the soils investigated the rate of corrosion of ferrous materials decreases as the exposure is prolonged. Any rate of corrosion is applicable only to the area of the metal tested and the time it was exposed. The life of a pipe cannot be predicted solely from the loss of weight or the depth of a pit at any given time. The corrosiveness of the soil can be indicated only by a formula which takes account of the characteristics of the soil to which the pine is exposed, the charge in the of the characteristics of the soil to which the pipe is exposed, the change in the rate of corrosion with time and the area of the exposed metal.

The data in the Circular show that copper and alloys high in copper corrode much less and more uniformly than ferrous materials. Lead was resistant to many soils

but pits deeply in some soils.

The effectiveness of metallic coatings depends on the soils to which they are exposed. No metallic coating is suitable for all soils. Bituminous coatings greatly retard the loss of metal to which they are applied. Their effectiveness is roughly proportional to their thickness. Few coatings are free from all pinholes and other imperfections. Important causes of coating failures are improper application and injuries incidental to pipe laying. Better tools and methods have considerably reduced the number of failures attributable to these causes, but it is still difficult to see the sould be soil stress. to secure a completely protected pipe and some coatings are injured by soil stress and roots of trees, shrubs, and grass.

The results of tests of a number of promising coatings developed in the laboratory

Several methods of testing soils and coatings are described and their usefulness discussed. The effectiveness of several methods of preventing corrosion are compared. Cathodic protection can retard or prevent corrosion under most soil conditions. The paper describes methods of applying cathodic protection under several conditions. Details of test methods and apparatus are given in six appendices.

CONTENTS

Page

	Prefac		Ι
I.	Introd	uction	7
II.		gations of underground corrosion	8
	11110301	In foreign countries	į
	2.	In the United States	9

III.	Causes	of underground corrosion	1(
	1.	General fromy	1(
	2.	Specific causes	12
	3.		13
		(a) Classification	13
		(b) Chemical properties	1.8
		(c) Physical properties	21
IV.	Turronti	gations by the National Bureau of Standards	24
т .			24
	1.	General plan and methods	
	2.		29
	3.	Soils at test sites	15
	4.	Factors affecting results of corrosion tests 5	54
		(a) Depth of burial 5	54
		(b) Mill and foundry scale 5	6
		(c) Diameter of pipe	56
			18
		(d) Area of pipe surface 5	
			1
v.	Results	of field tests on ferrous materials 6	2
	1.	Cast iron 6	2
	_,	(a) After 17 years of exposure	2
		(b) After 12 years of exposure	6
		(b) After 0 years of exposure	8
			3
			3
		(f) Malleable cast iron 7	3
	2.	Wrought ferrous material 7	4
		(a) After 17 years of exposure 7	4
		(b) After 12 years of exposure 7	
		(c) After 9 years of exposure	
	3.		
	4.	Bolts 9	
	5.		
VI.	Interpre	etation of results on ferrous materials 9	9
		General considerations 9	9
	2.	Extrapolation of data	0
	3.	Life of nine	5
	4.	Life of pipe	G
~~~~	T. 11	Comparison of test data with data on performance	7
VII.	Results	of field tests on miscellaneous nonferrous materials	T
	1.	Copper and copper alloys. 11 Lead 12	1
	2.	Lead	2
	3.	Zinc and aluminum 12	5
	4.	Asbestos-cement pipe	8
		(a) Water-absorption test	ñ
		(b) Crushing test	ň
		(a) Durating test	1
		(c) Bursting test         13           (d) Apparent specific gravity         13           (e) Comparison of data from various tests         13	T
		(d) Apparent specific gravity	3
		(e) Comparison of data from various tests 133	3
	5.	Parkway cables	4
TII.	Results	of field tests on protective coatings	4.
		Metallic coatings	
	-•	(a) Zine	i
		(b) Lead	~
			U
		(c) Aluminum (calorized)	4
		(d) Tinned copper 149	5
	2.	Bituminous coatings 14( (a) Scope and methods 14(	6
		(a) Scope and methods	6
		(b) American Gas Association tests	7
		(c) American Petroleum Institute tests	'n
		(c) American Petroleum Institute tests       140         (1) Sites and coatings       140         (2) Coatings applied to pipe lines       150	7
		(2) Continue and coatings	,
		(2) Coatings applied to pipe lines 153	L

		Page
	(3) Coatings applied to isolated short sections	1 70
	of small-dian acr pipe	198
	(4) Comparison of results of tests of coatings	
	applied to operating lines with those on	164
	isolated small pipes	-01
	coated pipe	168
	(6) Conclusions based on the API tests	176
	3. Nonbituminous, nonmetallic coatings	178
	(a) After 9 years of exposure	178
	(a) After 9 years of exposure	179
	(c) After 2 years of exposure	180
IX.	Laboratory studies of corrosion	181
	Denison corrosion cell     Behavior of steel in various soils	181
	2. Benavior of steel in various soils.	194
	3. Behavior of different metals in various soils	
X.		194
22.	Soil surveys and tests	194
	2. Field tests	195
	(a) Resistivity	
	(1) Shepard's soil-resistivity apparatus	196
	(2) Wenner's four-terminal method	197
	(3) Electrolytic-bridge method	197
	(4) Radio balance	198
	(b) Columbia rod	198
	(c) Redox apparatus	199
	(d) Identification of soil types	199
	3. Laboratory tests	208
	(a) Resistivity (b) Hydrogen-ion concentration and total acidity (c) Putnam's decomposition potential tests	219
	(c) Putnam's decomposition potential tests	214
	(d) Williams-Corfield or nipple-and-can test	215
	(e) Denison cell test	216
	4. Value of soil-corrosivity tests	216
	(a) Correlation of soil types with pit depths	218
	(b) Correlation of corrosion with field resistivity	219
	(1) Shepard's soil-resistivity measurements	219
	(c) Correlation of pit depths with soil resistivity at	
		$\frac{224}{224}$
	<ul><li>(d) Correlation of other soil test data with pit depths</li><li>(e) Relative merits of tests for soil corrosivity</li></ul>	226
XI.	Determination of the condition of a nine line	229
	Determination of the condition of a pipe line	$\frac{229}{229}$
	(a) Measurements of potentials and currents	229
	(b) Coupons	230
	(b) Coupons	
	pipe line	231
XII.		236
	1. Metallic coatings	236
	Insulating coatings	236
XIII.	Prevention of corrosion	241
	1. Materials suitable for use in mildly corrosive soils	241
	2. Corrosion-resistant materials	243
	3. Soil treatment	945
	(a) Concrete	245
	(b) Grease coatings	246
	(e) Bituminous coatings	246
XIV.	Cathodic protection	
	1. General principles	251
	2. Criteria of protection	251
	(a) Current density	251
	(b) Pipe-to-soil potential	252
	(c) Current-potential curves	254

		age
(d) Comparison of results of di	ifferent methods for de-	
termining the protective of		258
3. Cathodic protection installations		261
(a) Protection of transmission p	orpe imes	181
(1) Principles	t 2	669
(a) Converted	d power 2	62
(b) Primary	power 2	62
A. A	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	62
В. 1	Magnesium anodes 2	62
C. Z	line anodes 2	63
(3) Anodes		64
(a) Resistance	e to earth	64
		66
A. 1	ron 2	66
В. С		68
		69 69
(b) Cathodic protection of city of tank		69
(c) Cathodic protection of tank (d) Cost of cathodic protection.		70
(e) Status of cathodic protection		71
XV. References	2'	71
XVI. Appendix 1. Cooperators with the National		
the corrosion investigations	<i>.</i>	79
1. Furnishers of labor and of test sites		79
2. Suppliers of materials	28	80
(a) Ferrous		80
(b) Nonierrous		80
(c) Coatings		80
3. Advisory organizations		81 81
4. Government departmentsXVII. Appendix 2. Description of soils at test sites.	ند	оı 82
XVIII. Appendix 3. Methods employed in cleaning of	the National Rureau of	04
Standard soil-corrosion specimens		02
<ol> <li>Cleaning of iron and steel specimens.</li> <li>Cleaning of copper, copper alloy, and a</li> <li>Cleaning of lead and lead-coated speci</li> </ol>		02
2. Cleaning of copper, copper-alloy, and a	aluminum specimens 30	03
<ol> <li>Cleaning of lead and lead-coated speci</li> </ol>	mens 30	03
4. Cleaning zinc and galvanized specimen	18 30	04
5. Cleaning of calorized pipe		05
6. Cleaning of nonmetallic-coated specime	ens 30	05
XIX. Appendix 4. Accumulation of data, calculation	ns, and methods of re-	^-
porting the data		05
1. Accumulation of data		05 06
2. Calculations and methods of reporting XX. Appendix 5. Construction of the Denison corr XXI. Appendix 6. Darnielle's modification of the	ogion cell	07
XXI. Appendix 6. Darnielle's modification of the	Hickling apparatus for	٠.
measuring potentials	30	09
XXII. Index		11
TABLES		
<ol> <li>Description of typical profiles in great soil group soils in which the National Bureau of Standards</li> </ol>	ps, and classification of	15
2. Classification of soil particles as to size	s dest altes alteredated	$\frac{15}{16}$
3. Composition of water extracts of some corrosive an	nd noncorrosive soils	20
4. Relation of acidity to pipe line repairs in differen	t types of soils 2	21
5. Correlation of soil corrosivity with the physical pr	coperties of soils 2	25
5. Correlation of soil corrosivity with the physical pr 6. Maximum and minimum values of certain proper	rties of the soils at the	
National Bureau of Standards test sites		25
7. Composition of wrought ferrous materials		30
8. Composition of cast ferrous materials		32
9. Composition of copper and copper-alloy materials 10. Composition of zinc. lead, and aluminum-alloy materials	3	33 34
11. Metallic-coated specimens	ateriais	35
12. Nonbituminous-nonmetallic coatings		36
11. Metallic-coated specimens	al Bureau of Standards	, ,
tests		37
tests  14. Coatings in American Petroleum Institute line te	sts.,,,	38

		Page
15.		39
16.	Characteristics of bituminous coating materials in the American Petroleum	41
17.	Institute tests	42
18.	Characteristics of the American Gas Association coating materials	44
19.	Test sites, soil numbers, types, locations, and other information relating	
	to soils at NBS, AGA, and API test sites	<b>4</b> 6
20.	Chemical properties of soils at National Bureau of Standards test sites	50
21.	Physical properties of soils at National Bureau of Standards test sites	52
22.	Mechanical analysis of soils at National Bureau of Standards test sites	53
23.	Properties of soils in the Amercian Gas Association and American Petro-	=1
~ .	leum Institute coating tests	54
24.	Effect of depth of burial on corrosion of 3-inch steel pipe	55 56
25. 26.	Effect of mill and foundry scale on corrosion	50
۵0.	neriods of exposure	57
27.	periods of exposure	0.
~	Bureau of Standards specimens	59
28.	Relation between area inspected and the observed depth of the deepest pit	59
29.	Loss of weight and maximum and weighted maximum penetration of 17-	
	year-old cast-iron specimens	63
30.	Loss of weight, maximum, weighted maximum and estimated average	
0.7	year-old cast-iron specimens.  Loss of weight, maximum, weighted maximum and estimated average maximum penetration of 12-year-old cast-iron pipe.  Loss of weight and depth of maximum penetration of cast-iron pipe ex-	64
31.	Loss of weight and depth of maximum penetration of cast-fron pipe ex-	69
32.	posed for 9 years	U a
J2.	periods of exposure	70
33.	Average of the relative maximum penetration of the cast materials for	••
	four periods of exposure	71
34.	four periods of exposureLoss of weight and depth of maximum penetration of cast-iron plates and	
	cast-iron and steel bolts	73
35.	Corrosion of malleable cast iron and cast steel	74
36.	Loss of weight of 17-year-old wrought ferrous pipe	75
37.	Maximum penetration and weighted maximum penetration of 17-year-old	76
38.	wrought ferrous pipeLoss of weight of 12-year-old wrought ferrous pipe	80
39.	Maximum, weighted maximum, and estimated average maximum penetra-	00
	tion of 1½-inch wrought ferrous pipe exposed for 12 years	82
40.	Maximum, weighted maximum, and estimated average maximum penetra-	
	tion of 3-inch wrought ferrous pipe exposed for 12 years	84
41.	Loss of weight of wrought pipe exposed for 9 years  Depths of maximum penetration of wrought pipe exposed for 9 years	86
42.	Depths of maximum penetration of wrought pipe exposed for 9 years	87
43.	Average of the relative loss of weight of wrought specimens for four	88
44.	periods of exposure	00
	four periods of exposure.	88
45.	four periods of exposure	
	sheets exposed for 9 years	92
46.	Loss of weight of alloy iron and steel plates exposed for 4 years	93
47.	Maximum penetration of alloy iron and steel plates exposed for 4 years Relative loss of weight and maximum penetration of alloy iron and steel	94
48.	Relative loss of weight and maximum penetration of alloy iron and steel	0-
<b>4</b> 9.	plates	95
40.	allow steel specimens exposed for 2 years	97
50.	Corrosion of 3/-inch bolts	98
51.	Calculated data on pit depths and leaks on a pipe line	102
52.	Depth of the deepest pit on 4 feet of uncoated line pipe and on the	
	adjacent 3-inch control pipe.  Composition of cast brass caps.	110
53.	Composition of east brass caps	111
54.	Corrosion of copper-rich alloys exposed for 8 years and for 13 years	116
55.		118
56. 57.		119
58.	Loss of weight and maximum penetration of Muntz metal exposed for	120
٠	2 years	121
59.	Loss of weight and maximum penetration of lead cable sheaths exposed	
	10 to 17 years,	123

	Page
(d) Comparison of results of different methods for de-	
termining the protective current	258
3. Cathodic protection installations	261
(a) Protection of transmission pipe lines	261
(1) Principles	201
(2) Sources of current	
(a) Converted power	
(b) Primary power	
B. Magnesium anodes C. Zinc anodes	
(3) Anodes	
(b) Materials	
B. Carbon and graphite	
C. Noncorrodible materials.	
(b) Cathodic protection of city networks	
(d) Cost of cathodic protection	270
(e) Status of cathodic protection engineering	271
7777 10 - 0	
XVI. Appendix 1. Cooperators with the National Bureau of Standards in	
the corresion investigations	279
the corrosion investigations	279
2. Suppliers of materials	280
(a) Ferrous	280
(b) Nonferrous	280
(c) Coatings	280
3. Advisory organizations	281
4. Government departments	281
XVII. Appendix 2. Description of soils at test sites	282
XVIII. Appendix 3. Methods employed in cleaning of the National Burcau of	
Standard soil-corrosion specimens	302
1. Cleaning of iron and steel specimens	302
2. Cleaning of copper, copper-alloy, and aluminum specimens	303
3. Cleaning of lead and lead-coated specimens	303
4. Cleaning zinc and galvanized specimens	304
5. Cleaning of calorized pipe	305
6. Cleaning of nonmetallic-coated specimens	305
XIX. Appendix 4. Accumulation of data, calculations, and methods of re-	
porting the data	305
1. Accumulation of data	
2. Calculations and methods of reporting the data	
XX. Appendix 5. Construction of the Denison corrosion cell	307
XXI. Appendix 6. Darnielle's modification of the Hickling apparatus for	000
measuring potentials	309
XXII. Index	311
TABLES	
1. Description of typical profiles in great soil groups, and classification of	
soils in which the National Bureau of Standards test sites are located	15
2. Classification of soil particles as to size	16
3. Composition of water extracts of some corrosive and noncorrosive soils	20
4. Relation of acidity to pipe line repairs in different types of soils	21
5. Correlation of soil corrosivity with the physical properties of soils	25

## Underground Corrosion

15.	Coatings on American Petroleum Institute short pipe sections
16.	Characteristics of bituminous coating materials in the American Petroleu
	Institute tests
17.	Descriptions of pipe coatings in the American Gas Association tests
18.	Characteristics of the American Gas Association coating materials
19.	Test sites, soil numbers, types, locations, and other information relating to soils at NBS, AGA, and API test sites
20.	Chemical properties of soils at National Bureau of Standards test sites
21.	Physical properties of soils at National Bureau of Standards test sites.
22.	Mechanical analysis of soils at National Bureau of Standards test sites
23.	Properties of soils in the Amercian Gas Association and American Petr
	leum Institute coating tests
24.	Effect of depth of burial on corrosion of 3-inch steel pipe
25.	Effect of mill and foundry scale on corrosion
26.	Average of maximum pit depths on pipes of different diameters for for
07	periods of exposure
27.	Bureau of Standards specimens
28.	Relation between area inspected and the observed depth of the deepest p
29.	Loss of weight and maximum and weighted maximum penetration of 1
	year-old cast-iron specimens
30.	Loss of weight, maximum, weighted maximum and estimated average
	maximum penetration of 12-year-old cast-iron pipe
31.	maximum penetration of 12-year-old cast-iron pipe
	posed for 9 years
32.	Average of the relative loss of weight of the cast materials for for
	periods of exposure
33.	Average of the relative maximum penetration of the cast materials is
9.4	four periods of exposure
34.	cast-iron and steel bolts
35.	Corrosion of malleable cast iron and cast steel
36.	Loss of weight of 17-year-old wrought ferrous pipe
37.	Maximum penetration and weighted maximum penetration of 17-year-ol
	wrought ferrous pipe
38.	Loss of weight of 12-year-old wrought ferrous pipe
39.	Maximum, weighted maximum, and estimated average maximum penetra
	tion of 1½-inch wrought ferrous pipe exposed for 12 years
40.	Maximum, weighted maximum, and estimated average maximum penetr
47	tion of 3-inch wrought ferrous pipe exposed for 12 years
41. 42.	Loss of weight of wrought pipe exposed for 9 years
43.	Depths of maximum penetration of wrought pipe exposed for 9 years Average of the relative loss of weight of wrought specimens for for
10.	neriods of exposure
44.	periods of exposure
	four periods of exposure
45.	four periods of exposure
	sheets exposed for 9 years
<b>4</b> 6.	Loss of weight of alloy iron and steel plates exposed for 4 years
47.	Maximum penetration of alloy iron and steel plates exposed for 4 years.
48.	Relative loss of weight and maximum penetration of alloy iron and ste
<b>4</b> 9.	Loss of weight and depth of maximum penetration of low-alloy and high
	alloy steel specimens exposed for 2 years
50.	Corrosion of %-inch bolts
51.	Calculated data on pit depths and leaks on a pipe line
52.	Depth of the deepest pit on 4 feet of uncoated line pine and on the

## Circulars of the National Bureau of Standards

		T 448
60.	Loss of weight and maximum penetration of lead pipe exposed for 4 years	128
61.	Loss of weight and maximum penetration of leat pape exposed for 4 years  Loss of weight and maximum penetration of zinc plates exposed for 4 years  Loss of weight and maximum penetration of zinc and aluminum exposed	127
62.		
63.	Absorption of water by asbestos-cement pipe	129
64.	Results of crushing tests on the asbestos-cement specimens exposed to	
	Results of crushing tests on the asbestos-cement specimens exposed to various soils and on the unexposed specimens	131
65.	Results of bursting tests on the aspestos-cement specimens exposed to the	132
66.	Condition of parkway cable exposed 10 to 10 years	138
67.	Condition of parkway cable exposed 10 to 16 years	700
	2-ounce coatings	136
	Loss of weight and depth of maximum penetration of galvanized and	
	2-ounce coatings  Loss of weight and depth of maximum penetration of galvanized and black iron pipe exposed for 4 years.  Loss of weight and depth of maximum penetration of 1½-inch lead-	132
69.	Loss of weight and depth of maximum penetration of 1½-inch lead-	
		140
70.	Those of weight and depth of maximum penetration of 1-1/2-inch lead-	
	coated pipe and Bessemer steel pipe exposed for 10 years Loss of weight and depth of maximum penetration of lead-coated and	141
71.	Loss of weight and depth of maximum penetration of lead-coated and	
	steel pipe exposed for 9 years	142
72.	Corrosion of calorized steel pipe exposed for approximately 10 years	143
73.	Loss of weight and depth of maximum penetration of tinned-copper tubes	
	exposed for 4 years and copper pipe exposed for 5 years	145
74.	Averages of maximum pit depths on coated and corresponding bare pipes	
	with soil resistivities	152
75.	Summary of conditions of line pipe undercoatings	158
76.	Single deepest pit on 20 feet of coated line pipe after 10 years of	
	exposure	159
77.	Depth of the deepest pit on short pipe sections	160
78.	Summary of conditions of 3-inch coated pipe	162
79.	Summary of conditions of 3-inch coated pipe	
	isolated sections of small-diameter pipe.	164
80.	isolated sections of small-diameter pipe	168
81.	Maximum conductance of coatings on operating pipe lines	170
82.	Maximum conductance of coatings on operating pipe lines	178
83.	Condition of metal beneath Bakelite and Koroseal coatings exposed for	
	4 Vears	179
84.	Summary of data on condition of coated pipe exposed for 4 years	180
85.	Condition of coated pipe exposed for 2 years	180
86.	Corrosiveness of soils as indicated by electrical measurements and loss of	
	weight of the anodes	187
87.	Control of the corrosion rate of metals in different soil environments	190
88.	Comparison of corrosion data from laboratory and field tests	192
89.	Repairs to pipe lines in different types of soils as related to their resistivity	195
90.	Summary of data on soils and pipe-line repairs	201
91.	Repair data for Lordstown sandy loam	202
92.	Summary of data on soils and pipe-line repairs	
	eastern Unio and their corrosiveness	204
93.	Relation between the degree of development of the glacial soils of north-	
	western Ohio and their corrosiveness	204
94.	Relation of soil series to rates of loss of weight and to weighted rates of	
	maximum penetration of ferrous specimens	206
95.	Repairs in soil types as related to total acidity	213
96.	Corfield corrosivity index	215
97.	Corfield corrosivity index	
	depths on the joint of the specimen adjacent to the soil sample	227
	Comparison of methods of testing soils on basis of correct arrangement	
	of soils in the rank, or order, of corrosiveness	228
99.	Character of pipe-line fabric materials and results of breaking tests.	940
100.	Effect of cathodic protection on rate of leak occurrence.  National Burkan of Standards test sites.	261
101.	National Bureau of Standards test sites	282
102.	Enter of medianical deaning of the specimens	303
103.	Effect of oxalic acid cleaning solution on nonferrous specimens	303
104.	Effect of nitric and oxalic acids on lead specimens	304
105.	Effect of 10-percent ammonia solution on zinc specimens	304
106.	Loss of weight of zinc specimens caused by ammonia chloride bath and	
	scrubbing	304

### I. INTRODUCTION

The prevention of damage to underground structures arising from corrosion has long been an engineering problem of major importance. There are in the United States about 500,000 miles of pipe lines used for transporting water, gas, oil, and gasoline. The value of these lines is in the order of 6 billion dollars. Accurate figures for the corrosion losses of these lines are not available.

The average life of steel mains has been taken as 33½ years by the Interstate Commerce Commission.¹ On this basis, the annual cost for replacement of pipe lines would be 200 million dollars. Most of the loss is probably due to corrosion.

In 1910 Congress first made funds available for an investigation by the National Bureau of Standards of the causes of stray-current electrolysis and its mitigation. For 11 years the Bureau conducted field and laboratory tests on this subject. The publications dealing with the results of these investigations are out of print, but they are summarized in Circular C401. These investigations indicated that although very serious corrosion resulted from stray currents, similar corrosion occurred under circumstances that precluded stray currents as an explana-The Bureau, therefore, undertook to determine whether soils caused corrosion of pipe materials, and, finding some corrosive soils, began the study of the relation of the various properties of soils to corrosion and later of methods for reducing corrosion losses. In this work it had the cooperation of a large number of manufacturers of materials used underground and of the users of these materials. These organizations not only furnished materials and labor but took an active part in the formulation of the conclusions, based on the results of the tests. As the work progressed, the results and conclusions were published from time to time in the National Bureau of Standards Journal of Research and in technical and trade journals. Most of these publications are now out of print. Although these are on file in many public, technical, and college libraries, the data are scattered through so many publications that the results of the investigations are not conveniently available. It is desirable therefore to summarize the work on underground corrosion and to present the more important data in one publication.

There are more than 6,000 specimens of pipe materials still buried at the test sites, including some types on which no data have been obtained. There are also a number of corrosion phenomena that should be studied further. However, the work on underground corrosion has been suspended and the staff assigned to more urgent work for the duration of the war. Because of the need for the data already obtained, the results of the studies of underground corrosion are summarized in this Circular.

Although in many cases the data are insufficient to justify positive conclusions, nevertheless, it seems desirable to suggest tentative conclusions, with the idea of stimulating discussion and further research.

¹ Valuation Docket 1203, Interstate Commerce Commission (1937).

To the conclusions resulting from the investigations of the National Bureau of Standards have been added the results of the more important investigations made elsewhere, and a sufficient number of references have been given to enable the reader to familiarize himself with the details of these investigations insofar as they have been published.

# II. INVESTIGATIONS OF UNDERGROUND CORROSION 1. IN FOREIGN COUNTRIES

Although the most extensive investigations of underground corrosion appear to have been carried out in the United States, the records show that similar investigations have been undertaken in several other countries.

The Dutch Research Institute for Metals in 1931 set up a Central Corrosion Committee. This committee delegated its work to various subcommittees, which investigated, among other things the following topics: Corrosive effects of soils on pipes, corrosive effect of soils on cable sheaths, and protective coatings for steel construction. The subcommittee for the effect of soil on pipe consisted of 2 representatives of the Central Committee and 12 officials, engineers, and chemists of laboratories and oil distributors. After a study of foreign literature on corrosion, pipe lines were inspected in various parts of Holland, and laboratory studies of soils, soil waters, and pipe materials were made. The committee issued a report [1]² in 1935. This will be referred to later. A similar subcommittee on coatings issued a report [2] on coatings, and in 1937 a set of specifications [3] for the coating of pipes with asphalt. In addition to the reports of the committee, individual members of the subcommittees have published a number of papers [4, 5] resulting from their researches. Paralleling some of these papers is a series of reports by Wichers [6, 7].

In Australia several organizations have conducted extensive investigations of underground corrosion and protective coatings, as well as of stray-current electrolysis, the results of which were presented in the technical press and at the 1937 and 1943 National Bureau of Standards Corrosion Conferences [8 to 15]. Most of the conference papers have

not been published.

In England the Institution of Civil Engineers, the Iron and Steel Institute, and the British Nonferrous Metals Association organized a Subcommittee on Soil Corrosion of Metals, which, after an investigation and some preliminary experiments, issued an interim report [16] in 1942. A detailed study of anaerobic corrosion [17] has been undertaken in England also. The Post Office Engineering Department [18] has made studies of the corrosion of lead cable sheath and has developed special apparatus for this work. The British Board of Trade [19], after an investigation of stray-current electrolysis, issued a set of regulations to govern tramway installations in 1912, and these have been revised from time to time. The Chemical Research Laboratory of the Department of Scientific and Industrial Research, Teddington, and the Metallurgical Laboratories of Cambridge University have conducted very extensive investigations of corrosion processes of metals and alloys exposed to a large variety of gases and solutions and have done much to establish the electrolytic theory of corrosion. The results of these investigations have been summarized by Evans [20].

² Figures in brackets indicate the literature references at the end of this paper.

The study of corrosion has been very extensive in Germany. Straycurrent electrolysis has been regulated since 1910. The potential gradient in the rails and the earth were limited and pipe drainage was prohibited. In 1929 the first volume of what was to be a set of books on corrosion [21] was published. This book contained 53 pages of references to articles on corrosion and related subjects, most of which were published in Germany.

Corrosion, especially stray-current electrolysis, has been studied extensively in Japan. An engineer from the Research Department of the Japanese Railways spent a month at the National Bureau of Standards in 1932 studying corrosion literature and methods of testing, and later visited many of the laboratories studying corrosion. Japan contributed four papers to the 1937 Underground Corrosion Conference and was represented at the conference by three engineers.

Severe corrosion of water mains in Winnipeg, Manitoba, led to an extended study of corrosion at the University of Manitoba in Canada [22 to 27], which added considerable information on the corrosion of cast iron.

### 2. IN THE UNITED STATES

In the United States, underground corrosion has been recognized as a serious problem for many years, and most of the larger companies operating pipe lines employ from one to a dozen or more men whose chief duty is to reduce pipe-line corrosion. One large gas company collected a sample of soil at each point where one of its pipes was exposed until soil conditions through its territory were determined. The data have not been published, but several papers based on them have appeared. Another large gas company has made a statistical study of the service pipe that it has installed at various times. Many pipe-line operators have experimented extensively in the laboratory and in the field with a large variety of protective coatings. From these activities a large quantity of data has accumulated which as yet has not been adequately studied, although many papers based on the data have been published. In many cases definite conclusions cannot be drawn from the corrosion records because the conditions under which the corrosion occurred are not described in sufficient detail. This is especially true with respect to soil conditions.

In 1928 or 1929 the American Gas Association and the American Petroleum Institute each formed a committee on corrosion and employed Research Associates to work at and under the direction of the National Bureau of Standards. For many years thereafter the annual proceedings of these associations contained articles on corrosion by these men and other members of the Associations. Many additional articles on corrosion appeared in technical and trade journals during

this period.

One of the most extensive and long-continued studies of corrosion is the investigation by the National Bureau of Standards previously referred to. The data from this investigation have been widely quoted and many conclusions have been drawn therefrom. It should be borne in mind, however, that the tests were planned with definite objectives in view and that, although the data were satisfactory from the point of view of these objectives, it will be necessary to guard against attempting to draw more far-reaching conclusions from them than are justified by the conditions under which the tests were made. In view of the widespread use of the National Bureau of Standards

corrosion data, the tests will be described and the interpretation of the results discussed in considerable detail.

The seriousness of underground corrosion was recognized by Congress in 1910, when the National Bureau of Standards was authorized to make a study of electrolysis caused by stray electric currents and possible methods of its mitigation. The work was carried on actively during a period of 10 years, and the results are reported in 17 Technologic Papers of the National Bureau of Standards. These papers discuss various phases of electrolysis, including surveys, track leakage, electrolysis in concrete, protective coatings, and the design of return feeder systems for street railways. Practically all these papers are now out of print and available only in reference libraries, but abstracts of the papers were published in 1933.

The work of the National Bureau of Standards on corrosion due to causes other than stray currents was started in 1922 and was actively carried on until interrupted by the necessity of giving full attention to war work. In this work, the Bureau has had the support and cooperation of a large number of makers and users of underground pipe and pipe coatings. These organizations have borne much of the expense of the investigations, and their technical representatives have had an active part in planning the tests and in formulating conclusions therefrom.

The results of these investigations, including studies of special phases of underground corrosion carried out by Research Associates working under the direction of the Bureau, have appeared in publications of the Bureau and in publications of the organizations supporting the research projects. In addition, a large number of papers by members of the Bureau staff have been presented before technical societies and published in technical journals.

There are so many diverse factors that affect the corrosion of pipes and other structures underground that the planning of adequate tests and the proper interpretation of the results are matters of considerable difficulty. It is not surprising, therefore, that even experienced corrosion engineers often interpret the same experimental data in different ways. For this reason, it is necessary to guard against the acceptance of too far-reaching or erroneous conclusions, which may be based upon insufficient data or which may be due to failure to take into consideration all the pertinent facts. Unfortunately, it is necessary also to warn against taking at face value the statements contained in publications obviously prepared for the purpose of promoting the sale of some particular product. On the other hand, it is necessary to make as much use as possible of the limited data because the cost of tests sufficiently extensive to warrant definite conclusions on a statistical basis would be prohibitive, even if such tests were feasible. In evaluating the conclusions presented in various publications, therefore, it is always well to take into consideration the experience and standing of their authors.

### III. CAUSES OF UNDERGROUND CORROSION

### 1. GENERA'L THEORY

Although the theory of corrosion in general is fairly simple and easily understood, the relative importance of the factors that affect corrosion underground is so difficult to evaluate and the conditions under which corrosion occurs are so poorly defined that the theory is often more help-

³ A list of the cooperating organizations is given in the Appendix 1.

ful in explaining corrosion that has already taken place than it is in predicting what may be expected. Nevertheless, it is essential as a guide in studying corrosion phenomena and in interpreting the experimental results.

Speller [28] in his book on corrosion gives an easily understood discussion of the electrolytic theory of corrosion. Ewing [29] in a handbook prepared for the American Gas Association, gives an abstract of Speller's discussion. Many phases of corrosion, both from the theoretical point of view and from the practical standpoint, are discussed by Burns and Schuh [30], Hoar [31], and Evans [20]. An extensive discussion of the origin of potential differences leading to corrosion is given by Gatty and Spooner [32]. The following summary of these references, although necessarily too brief to constitute a comprehensive treatment of the theory and phenomena of corrosion, may serve as a guide to the interpretation of the results of the investigations and tests given in this Circular and suggest ways in which corrosion may be at least partially prevented.

Most metals when exposed to air quickly become covered with a film of oxide, which tends to render them more noble, or electropositive, and less corrodible. Unfortunately, this oxide film is often either discontinuous or unstable. If the film-coated metal is in contact with an electrolyte, such as moisture condensed from the atmosphere or with water from some other source, metal particles bearing electric charges (cations) leave the surface of the metal at points where the film is discontinuous (anodes). The simultaneous emission of an equivalent number of electrons from the oxidized surface (cathode) results in the formation of hydroxyl ions, with consequent removal of hydrogen. This migration of ions constitutes a flow of current. The reactions which follow depend on the availability of oxygen and the chemical composition of the electrolyte. The reactions at the anodic and cathodic areas tend to change the potentials of the respective surfaces in opposite directions, thus causing the potential difference to decrease. The potential of the metal with respect to the adjacent electrolyte is a measure of its tendency to discharge ions, that is, to corrode, but this potential does not determine the rate of corrosion. The increase of the potential at the anode because of the accumulation of metal ions in solution and the decrease of the potential of the cathode by the deposit of hydrogen or other processes are known as anodic and cathodic polarization, respectively. It is this polarization of the electrodes that controls the rate of corrosion rather than the potentials recorded in the electromotive series.

Attention is called to the fact that the explanation of corrosion processes began with the assumption of an oxidized and an oxygen-free area on the surface of the metal, the oxidized surface being the one that did not corrode further and was at the higher, or more noble, potential. If this is true, it should be expected that if oxygen can reach some part of a metal surface and thus maintain the oxide film, the part of the surface less subject to oxidation will be anodic to the oxidized surface. This relation between well- and poorly-aerated surfaces has been demonstrated in the laboratory and is the cause of many cases of corrosion, especially in soils. Differences of potential may also occur when two metals are in contact, when the structure of the metal is not uniform because of strains or segregations of impurities, or when the metal contains two distinct materials, such as graphite and ferrite in cast iron. These conditions may be regarded as special cases of points on the

metallic surface that differ in solubility, solution pressure, or electric potential, which terms are different ways of expressing the same phenomenon. Similar conditions occur when the electrolyte in contact with different parts of the metal differs in density or composition.

In the papers referred to [30, 31, 32] the authors have discussed in detail corrosion processes resulting from the exposure of a wide variety

of metals and alloys to a large number of solutions.

Corrosion processes in soils are fundamentally the same as in other environments, but the study of underground corrosion is more difficult because the exact soil environment cannot be determined readily, and in many cases this environment is neither uniform nor stable. In brief, the corrosion process is as follows: The positive ions, which travel toward the cathode, may be discharged there. Frequently, however, they lose their charges en route. Under such conditions hydrogen ions are discharged at the cathode. The hydrogen ions may reduce some of the oxide film on the cathode or combine with oxygen in the electrolyte to form hydroxyl ions (alkali) or water or be deposited on the cathode as molecular hydrogen. The collection of alkali or hydrogen at the cathode makes the cathode more noble and reduces the corrosion current. This is one form of cathodic polarization. If ferrous ions leaving the anode contact OII ions in the solution, ferrous hydroxide is formed. Reaction with oxygen results in the formation of ferrous or ferric oxide (rust). If the rust is formed at the anode, it may reduce the rate of corrosion. If the combination occurs at a point remote from the anode, more ferrous ions enter the solution, that is, corrosion continues. Because of the many factors that influence corrosion in soils, it is usually impossible to predict accurately the performance of buried metal or to explain the cause of underground corrosion with assurance. However, although there is necessarily some uncertainty regarding specific cases of corrosion of buried metal, the results of a large number of tests and of observations of corroding pipe lines and other structures have made possible the prediction of the average performance of materials under specified conditions. A large amount of data on underground corrosion has been assembled, which, when generally understood, should aid materially in considering ways and means of reducing losses due to corrosion.

### 2. SPECIFIC CAUSES

The more important specific causes of underground corrosion may be listed in the order of their relative importance as follows:

Differential aeration.

Acids and salts in the soil.

Differences in characteristics of the soil at different points of contact. Foreign materials, such as cinders, scraps of metal, or organic materials.

Stray currents (in a few localities).

Nonuniformities in structure or conditions of the metal.

Use of material unsuitable for a given environment.

Interconnection of dissimilar metals.

Anaerobic bacteria.

Long-line currents.

Differential aeration generally results from unequal compactness of the backfill surrounding the buried metal, unequal porosity of the soil at different points, uneven distribution of moisture, and the restriction of the movement of air and moisture in the soil by buildings, roadways,

pavements, and vegetation. Acids and salts in the soil affect the tendency of metals to go into solution. Any differences in the characteristics of the soil in contact with points on the surface of the metal that effect the solubility of the iron lead to corrosion. Cinders, scraps of metal, and organic material may create differences of potential and thus cause corrosion. Stray currents from electric generators have been, in the past, an important cause of severe corrosion. This cause has diminished in importance during the last decade, but is still important in some localities. Differences of potential resulting from the way in which the metal was made are of only minor importance. Cases of segregation, improper heat treatment, poor or dirty material, etc., occasionally occur, but they are responsible for only a small percentage of losses due to corrosion. The use of materials unsuited to their environment is a much more frequent cause. The interconnection of different metals, such as the use of copper service pipe with iron water mains, occasionally results in corrosion. Anaerobic bacteria may be indirect causes of corrosion under certain conditions because their life processes depolarize cathodic areas. It is a question whether so-called long-line currents are the cause or the result of corrosion. The origin of these currents, which are frequently observed in cross-country pipe lines, is in doubt as they could result from one or more of several causes.

# 3. CHARACTERISTICS OF SOILS AS CORROSIVE ENVIRONMENT (a) CLASSIFICATION

The fact that the rate of corrosion and the depth and distribution of pits are different in different soils suggests the possibility that there may be a definite correlation between the type of soil and its corrosive action. A soil may be considered as a chemical that attacks a metal placed in it. However, the percentage of chemically active materials in most soils is very low and thus the solution is very weak. On the other hand, the chief factor in most underground corrosion is differential aeration. The distribution of air or oxygen and carbon dioxide in a soil, as well as the amount and distribution of moisture and ionized salts and acids, varies widely with the type of soil. For this reason, corrosion is more severe in some types of soils than in others.

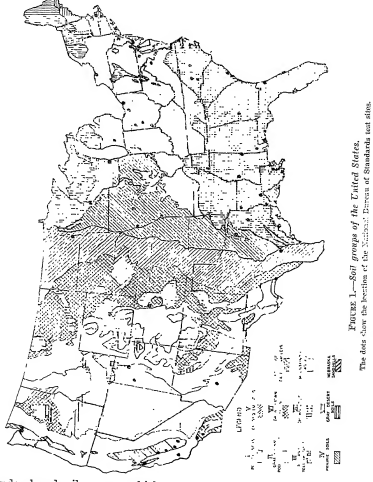
It should be pointed out that, in the strictly technical sense, the material in a trench in which a pipe is laid loses its identity as a soil when its structure is changed by the shovel or ditching machine. The structure of the backfill depends not only upon the soil in which the trench is located, but also upon the way in which the soil was handled. Nevertheless, the characteristics of the soil are important factors, and its classification can be very helpful in connection with the maintenance of pipe lines

A soil may be defined as two or more layers of finely divided, modified rock material having well-defined characteristics. In general, the term "soil" is applied to the first few feet of material covering the level and moderately inclined portions of the earth. Crushed rock and unmodified rock on mountain tops are not soils and, strictly speaking, earth removed from its original position is not soil if it has lost its structure.

The classification of soils is based on their physical and chemical characteristics and not on their geologic origin or geographic location, although the soil characteristics may be influenced by both the origin and the location of the soil.

Soils may be divided broadly into two classes: those in which lime

accumulates in the subsoil (pedocals) and those in which it does not (pedalfers). In the United States the first class lies generally west of a north and south line from northwestern Minnesota to a point on the Gulf of Mexico, 100 miles north of the Mexican border. Marbut [33] has classified the well-developed soils of the continental United States into eight great soil groups, to which he has added several groups of



undeveloped soils, among which are muck, peat, rough stony land, sand, and some others. Figure 1 shows the boundaries of the great soil groups, as defined by Marbut. Table 1 lists Marbut's great soil groups, briefly describes typical soil profiles, and gives some of the types represented

Table 1.—Description of typical profiles in great soil groups, and classification of soils in which the National Bureau of Standards test sites are located.

Class	Soil groups	Description of profile	Reaction of profile	Soil type
I	Podsol	A0, a few inches of leaf mat. A1, very thin dark-gray A2, whitish-gray, a few inches thick B1, dark- or coffee-brown B2, yellowish-brown	Strongly \acid	None.
I	Brown podsolic	(A0, leaf mat and acid humus	Acid	Gloucester sandy loam. Merrimac gravelly sandy loam.
II	Gray-brown podsolie	Ao, thin leaf litter and mild humus. Ai, dark-colored, 2 to 4 inches thick. Az grayish-brown, leached horizon extending to 8 to 10 inches. B, yellowish-brown to light red- dish-brown, definitely heav- ier in texture.	}do	Hagerstown loam. Chester loam. Sassafras silt loam. Miami clay loam. Lindley silt loam.
II	Yellow podsolic	As, thin, dark-colored organic covering. As, pale yellowish-gray leached layer Ginches to 3 feet thick. B, heavy, yellow. C, red and gray mottled parent material (acid).	}do	Kalmia fine sandy loam.  Norfolk sandy loam.  Ruston sandy loam.
111	Red podsolic	A6, thin organic layer	Acid	Cecil clay loam. Mem- phis silt loam. Sus- quehanna clay.
IV	Prairie	A, very dark brown or grayish-brown. B, brown C, light-colored parent material at 2 to 5 feet.	Slightly acid surface soil. Neutral to alka- line sub- soil.	Marshall silt loam. Summit silt loam.
v	Chernozem	A, black or very dark grayish- brown friable soil to a depth ranging from 3 to 4 feet B, light-colored to whitish lime accumulation	Alkaline	Fargo clay loam.
	∫Dark brown   soils	A, dark-brown or dark grayish- brown	do	
vII	Brown soils	A, brown	do	
	∫Gray desert { soils	A, light grayish-brown or gray, low in organic matter B, light in color, high in lime, often high in soluble salts	}do	(Mohave sandy loam. Panoche clay loam.
VIII	Soils of Pacific valleys	A, brown, reddish-brown or red, friable soil.  B, heavy, tough, more intense in color.	$\left\{egin{array}{l}  ext{Neutral} \\  ext{to} \\  ext{slightly} \\  ext{alka-} \\  ext{line} \end{array} ight\}$	Ramona loam.
VIII	Rendzina (im- mature soils developed from marl)	A, dark-gray or black granular soil B, soft, light-gray calcareous ma- terial.	Alkaline	{Houston black clay loam.
vIII	Alluvial soils			Genesee silt loam. Mil- ler clay. Sharkey clay. Wabash silt loam.
vIII	Saline soils	A, light-colored ashy material B, darker-colored heavy material of columnar structure	do	Docas clay. Fresno fine sandy loam.

in the National Bureau of Standards tests. Baldwin, Kellogg, and Thorp [34] in general followed Marbut's grouping of soils but have added several groups. It was not practicable at the time this Circular was prepared to classify all the soils in which test sites have been established by the Bureau.

According to the practice of the U.S. Department of Agriculture, the surface layer, or horizon, is designated by the letter A, the subsoil by B,

and the partly weathered parent material by C.

Each great soil group contains from 9 to 60 subgroups known as soil series, each of which is further subdivided as to the texture of the uppermost horizon. The three horizons comprising the profile of most soil types may differ widely in texture, color, and composition. A soil name consists of two parts—the first designates the series and the second the type. The series name is usually taken from the name of the locality where the series was discovered. The type name describes the texture of the top, or A, horizon. More than 260 soil series have been identified in the United States. Considerably more than two-thirds of the tillable soil of the United States has been mapped. Soil reports are issued for single counties or similar political subdivisions. Each report contains descriptions of the soil types and a map that shows in color the locations of the soil types within the area. The reports do not discuss the corrosivity of the soils, but they are, nevertheless, very helpful, as the pipe-line engineer can learn from them the aeration, drainage, and other characteristics of the soil that affect corrosion along his right-ofway. Unfortunately, most of the desert and mountain lands where oil and gas are frequently found have not been mapped. The soils of the older cities have not been mapped, but soil surveys of cities would be of little value because of the disturbing effects of cuts, fills, pavements, and buildings.

The texture of a soil horizon is determined by the percentages of the particles of various size groups. As to particle size, there are two grand subdivisions—those materials having diameters of 2 millimeters (0.079 inch) or more, which includes gravel, cobbles, and larger stone, and a group of materials of smaller diameter subdivided as indicated in table 2.

Table 2.—Classification of soil particles as to size.

Class	Diameter, millimeters
Gravel and stones. Fine gravel. Sand Silt Clay	1 to 2 0.05 to 1 0.002 to 0.05

Marbut has classified soils as to texture as follows:

- Sands include all soils containing 20 percent or less of silt and clay, the rest of the material being sand.
  - Coarse sands contain 35 percent or more of fine gravel and coarse sand and less than 50 percent of other grades of sand.
  - Medium sands contains 35 percent or more of fine gravel, coarse and medium sand, and less than 50 percent of other grades of sand.

⁴ Soil survey reports may be secured from the Superintendent of Documents, Government Printing Office, Washington 25, D. C. Give name of county and State.

- 3. Very fine sands contain 50 percent or more of very fine sand. II. Sandy loams contain from 20 to 50 percent of silt and clay. They are designated as coarse, medium, fine, and very fine sandy loams in accordance with the predominant sand class group present. There are also gravelly loams and stony loams.
- Loams and clays contain 50 percent or more of silt and clay combined.
  - 1. Loams contain 20 percent or less of clay, from 30 to 50 percent of silt, and from 30 to 50 percent of sand.
  - Silt loams contain 20 percent or less of clay, 50 percent or more of silt, and 30 percent or less of other classes.
  - 3. Clay loams contain from 20 to 30 percent of clay, from 20 to 50 percent of silt, and from 20 to 50 percent of sand.
  - Clays contain 30 percent or more of clay and 70 percent or less of other classes.

The classification of soils as to texture is shown in figure 2. This figure

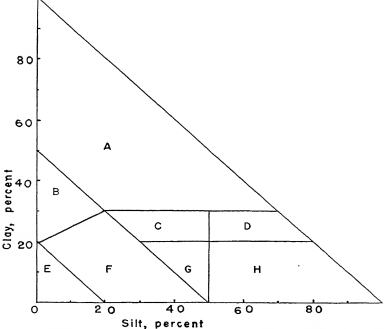


Figure 2.—Whitney diagram for classifying soils according to their clay and silt content.

The texture of the soils is determined by the area in which the clay and silt ordinates intersect:

A, clay; B, sandy clay; C, clay loam; D, silty clay loam; E, sand; F, sandy loam; G, loam;
H, silt loam.

shows only the percentages of clay and silt. Their sum subtracted from 100 gives the percentage of sand.

The A horizon, the texture of which determines the soil type, is usually less than a foot in thickness and may differ in many ways from the

horizon below it, in which pipes are commonly laid. Beneath the true soil are one or more layers of unconsolidated material from which the soil is being slowly formed. As pipes usually do not lie in the A horizon, the name of the soil type does not indicate the texture of the soil to which the pipe is exposed. However, as a soil with a given name is always substantially the same wherever it occurs, the name of the soil does disclose to those familiar with its significance the character of the soil in all its parts. Thus, wherever found, the light-textured A horizon of Susquehanna sandy loam is underlain by a heavy, impervious clav subsoil. The Norfolk sandy loam, on the other hand, always has a friable sandy clay subsoil. Consequently, by reading the soil descriptions which accompany the soil maps, the pipe-line engineer can determine whether a section of his line will be exposed to a poorly drained soil, where corrosion is likely to be bad and where soft coatings would be distorted, or to a well-drained soil, where little corrosion is to be anticipated. (b) CHEMICAL PROPERTIES

Although a very large number of chemical elements exist in soils, most of them are combined in difficultly soluble compounds, which exert little chemical influence on corrosion. These inert components of soils are chiefly combinations of oxygen with silicon, aluminum, and iron. Iron in various degrees of oxidation is responsible for the color of many soils, and this color is an indicator of the degree of aeration of the soil.

Chemical analyses of soils, from the standpoint of corrosion, are usually limited to determinations of the composition of the solution resulting from the contact of water and soil under standard conditions. The base-forming metals usually looked for are sodium, potassium, calcium, and magnesium. The acid radicals sought are carbonate, bicarbonate, chloride, nitrate, and sulfate. Usually the hydrogen-ion concentration of the solution, or the total acidity of the soil, or both, are determined.

The development of acidity in soils is a result of the natural processes of weathering under humid conditions. In regions of moderate rainfall, soluble salts do not accumulate except where soil waters seep to lower levels and collect in depressions. However, in regions of high rainfall, not only are soluble salts removed from the soil but also the absorbed bases normally in the colloidal materials of the soil are partially removed and replaced with hydrogen ions. This process eventually gives rise to the condition known as soil acidity. The depths to which this replacement of bases occurs varies with rainfall, drainage, type of vegetation, and nature of the material present. The fully developed soils of the prairie regions are acid to a considerable depth, whereas soils whose development has been retarded by poor drainage or other conditions are alkaline and may even contain appreciable quantities of salts. East of the prairies the well-developed soils are acid throughout the soil profile.

The soluble constituents of soils affect corrosion because they control the potentials of the anodes and cathodes and the resistance of the path of the corrosion current. The soluble constituents, especially the salts, furnish most of the ions which carry the current. As a rule, soils containing considerable quantities of salts in solution are corrosive.

Corrosion is also affected by the reactions between the soluble salts and the primary products of corrosion, the effect on the rate of corrosion depending on whether or not the reaction products are precipitated and on the location of the deposited products with respect to the anode

and cathode of the corrosion circuit. In general, if the reactions result in the formation of soluble products, or if the products are precipitated at a distance from the electrodes, corrosion continues.

If insoluble, poorly conducting products are deposited on one or both of the electrodes, corrosion will be retarded. For example, iron continues to corrode in soils containing sodium chloride because the anodic corrosion product, ferrous chloride, and the cathodic product are both soluble. On the other hand, if calcium bicarbonate is present, difficultly soluble calcium carbonate is precipitated at the cathodic areas because of the increase in alkalinity in that region resulting from the flow of current. As a result of this deposit the current is reduced and the rate of corrosion decreases. However, if salts forming soluble corrosion products are present, the beneficial effect of calcium carbonate is largely nullified.

If iron is placed in a soil containing a soluble sulfate, corrosion progresses as in the presence of sodium chloride, but if lead is substituted for iron, corrosion ceases after a short time because of a deposit of insoluble lead sulfate.

Table 3 shows the composition of the water extracts from five of the least corrosive soils in the original National Bureau of Standards soil-corrosion tests and from five of the most corrosive soils in the same tests. The table also shows the maximum pit depths on the 12-year-old Bessemer steel specimens exposed to these soils, and the electrical resistivity of the soils. Analyses were not made of soils of high electrical resistivity because of the very low concentration of soluble material in these soils.

The rate of corrosion is affected by the time of exposure and the area of the specimen exposed. The apparent relative merits of materials with respect to corrosion may change with the duration of the exposure because of the development or breakdown of corrosion products. Likewise, the apparent relative corrosiveness of two soils with respect to a material may change with the period of exposure because of the effect of the soils on the development of protective corrosion products or the exhaustion of corrosive elements in one of the soils. For this reason, table 3 also includes five of the least corrosive soils and five of the most corrosive soils, based on the estimated time required for 1,000 feet of 8-inch steel pipe having a wall thickness of 0.322 inch to be punctured by corrosion. The paper [35] from which the data were taken indicates that the estimates are very rough. It will be noted that the corrosive soils are considerably lower in resistivity and higher in salt content than the less corrosive soils.

The effect of acidity on the corrosion of ferrous metals in soils and in natural waters is much the same as that of salts, such as sodium chloride, as the effect in both cases is to increase the solubility of the corrosion products. Baylis [36] has pointed out that the tendency of iron to go into solution would be a function of the hydrogen-ion concentration if there were rapid displacement of the solution and no tendency to form a surface coating. Whitman, Russell, and Altieri [37] and Whitman and Russell [38] found that in a solution free from salts which form protective coatings, the corrosion rate of steel was unaffected by the hydrogen-ion concentration between pH 9.5 and pH 4, but was determined by the rate at which oxygen diffused to the cathodic surface and by the protectiveness of the film formed on the metal surface. The protective action of this film in water having a pH as low as 4 was attributed to the fact that the quantity of hydrogen ions

TABLE 3.—Composition of water extracts of some corrosive and noncorrosive soils."

[mg-eq =milligram equivalent]

	5.5	Corrosivity	ivity		Acid	Acidity	Con	ryosition	of water e	tract, m	Composition of water extract, mg-eq per 100 g of soil	ю в об во	
No.	Type	Maximum pit depth after	ife	Electrical resistivity	Hq	Total mg-eq per 100 g of soil	Na+K as Na	Ca	Mg	CO3	нсоз	D	*0s
	12 years   12 years   12 YEARS   12 YEARS   12 YEARS   12 YEARS   13 YEARS   14 YEARS   15 YEARS	12 years   H RESPEC	T TO MAZ	XIMUM PI	T ON	126 IN.2 (	OF BESSI	EMER S	TEEL AT	r 12 ye.	ARS		
47. 24. 26.	Unidentifie Merrimac Miami silt Everett gra	Mils 29 31 39 40 45	Years	Ohm-cm 1,770 11,400 2,980 45,000 20,500	7.6 7.3 5.9	3 0 12.6 2.6 12.8 1.8	0.67	0 72	0.39	00 0	0.88	0.06	0.48
31	_	I BESPECT	TO MAX	IMOM PIT	r ON 126	% IN.2 O	IN, OF BESSEMER STEEL	MER ST	EEL AT	AT 12 YEARS	RS		
23 29 45	Merced (a Montezun Muck	173 153 146 137		278 408 1,270 263 350	4.8.2.4.0	PA (©) A A A	8.38 1.50 2.15 8.15 1.42	0.38 .06 1.92 3.70 1.72	0.22 .18 1.55 0.70 2.55	.0 .0 .0 .0 .0 .0 .0 .0 .0	1.87 0.12 .00 .24 .71	1.12 0.99 1.69 0.18	6.57 0.89 2.30 11 98 4.43
80	Fargo clay loam   11.9	ESPECT TO	LIFE OF	1,000 FT	OF 8 II	N. STEE	STEEL PIPE (BASED ON	BASED (	ON EWIN	NG'S EQ	EWING'S EQUATION) ⁴	p -	-
31 6 24	Norfe Ever Merr Rust		do 9×10¢ 5×10⁴ 9×10³	20,500 45,000 11,400 11,200 5,9S0	4.70.4.4.4. 7.0.10.10.10	1.8 12.8 12.6 4.6 19.1							
17	ASPECT OF LOOP TITTE RESPECT TO LIFE OF 1,000 FT OF	PECT TO	LIFE OF 1	,000 FT OI	F 8 IN.	STEEL	STEEL PIPE (BASED ON	SED ON	EWING	EWING'S EQUATION)	ATION)		-
28 23 28 28 28 28 28 28 28 28 28 28 28 28 28	Men Mo Pea Pea		7 8 10 13 16	1,270 278 408 800 263	4.6 6.8 4.7 7.4.8 8.4.2	28.1 (b) 36.0 A	8.33 1.50 8.15 8.15	1 92 0.33 7.30 3.70	1.55 0.22 1.18 4.06 0.70	0000000	0 00 1 87 0.12	1.69 1.12 0.99 0.99 0.18	2.30 5.57 0.89 2.13 11.98
70	-												

a Analyses by I. A. Denison.

A indicates also had had no.

on by the determined.

See reference [35] for discussion of Ewing's equation.

coming in contact with the metal surface was insufficient to neutralize and dissolve the film. However, when carbon dioxide was introduced rapid corrosion occurred at pH 5.4 because then the increased total acidity of the water was sufficient to neutralize the hydroxide film. This result led to the conclusion that total acidity is more important in corrosion by natural waters than is the actual concentration of hydrogen ions, that is, the pH value.

As the acids in soils are only slightly dissociated, the pH value of a soil may offer no indication of the capacity of the acidic material to prevent the formation of otherwise protective hydroxide films. An acid soil having a relatively high pH value and a high total acidity would be expected to be more corrosive than a soil having a lower pH value and a lower total acidity because of the greater tendency of high total acidity to prevent the formation of protective films. Denison and Hobbs [39] found a correlations between total acidity and the corrosiveness of soils with respect to steel, both in the field and in the laboratory, under conditions in which other causes of corrosion were nearly constant. This correlation was confirmed by Denison and Ewing [40], who obtained the correlation shown in table 4 between total acidity and replacements for a pipe line in Ohio. It will be noted that in the absence of other variables, the corrosiveness of the soils increased with their acidity. However, the standard deviation of the average percentage of repairs shows that even within a single type of soil the corrosion varies greatly.

TABLE 4.—Relation of acidity to pipe line repairs in different types of soils.

Soil type	Total acidity	Standard error	Repairs	Standard deviation
Wauseon fine sandy loam ^a Canaedea silt loam Miami silt loam Mahoning silt loam Trumbull elay loam Crosby silt loam	12.7 16.8 18.1 21.1	Percent  2.1 2.8 0.7 2.3 1.2	Percent 6.3 13.3 22.8 20.9 20.0 30.8	Percent 11.0 30.6 13.3 20.0 16.9

^a One sample.

In certain regions of the Midwest and Pacific Coast a correlation has been found between soil resistivity and corrosion, the soils having the lowest resistivity being the most corrosive. As the conductivity of the soils is entirely electrolytic, the amount and kinds of salts in solution would largely determine the resistivity of the soil, and the corrosivity of the soil would increase with the salt content.

### (c) PHYSICAL PROPERTIES

The physical properties of soils that are of importance in corrosion are chiefly those which determine the aeration of the soil and its retentiveness for water. The texture of the soil, which is determined by the percentages of the particles of the various size groups, is obviously an important factor with respect both to aeration and to moisture content. In soils of coarse texture, such as sands and gravels, in which there is free circulation of air, corrosion approaches the atmospheric type. However, in heavy clay soils, which are usually very retentive of water, corrosion proceeds in an atmosphere very deficient in oxygen, thereby tending to approach the condition of submerged corrosion. There are, however, differences between corrosion processes in water and in soil. Some of these occur because soils are much less homo-

geneous than water. Other differences occur because corrosion products diffuse more slowly in soils, as convection currents and other movements of water are retarded by soil particles. Moreover, corrosion tests in water usually involve a single corrosive agent, whereas in soils several different factors influence corrosion and may react on each other. For these reasons great care should be exercised in applying the results of tests in water to corrosion in soils.

Corrosion is retarded by the oxidation of corrosion products that produce films or thicker deposits. These deposits either reduce the differences of potentials between anodic and cathodic areas or interpose electric resistance which reduces the corrosion current. The degree to which oxidation occurs depends largely on how readily oxygen can reach the corroding metal, which is determined by the permeability of the different soil horizons both above and below the buried metal and the rainfall and drainage of the region. In general, the aeration of clay soils is poor, whereas that of sandy soils is usually good. However, low-lying sands, and sands underlain by a consolidated stratum or a hard-pan layer, may be poorly aerated for long periods in regions where the rainfall is heavy.

As the aeration of the soil, which is the chief factor in the supply of oxygen and carbon dioxide, is determined not only by the character of the soil but also by the amount of water that the soil contains, the aeration will vary from season to season and cannot be expressed exactly or numerically, except perhaps for some temporary condition.

In well-aerated soils the rate of pitting of ferrous metals is initially very high because of the abundant oxygen supply at the cathodic areas. However, oxidation and consequent precipitation of the corrosion products in close contact with the anodic areas cause a marked reduction in the rate of corrosion with the result that the ultimate depth of pitting is relatively slight. On the other hand, in poorly aerated soils the rate of pitting, although low because of deficiency of oxygen at the cathodic areas, is relatively unchanged with time, because the corrosion products in the reduced condition are precipitated at points remote from the anodic areas. Consequently, the depths of the deepest pits after a long period are usually considerably greater in poorly aerated soils than in well-aerated soils.

Some soil water occupies the larger spaces between the soil particles, holes left by decaying roots, etc. This part of the soil moisture varies greatly with the season, rainfall, and drainage. Water also adheres closely to the minute soil particles, and is removed from them and the small soil capillaries only with difficulty. On this account, well-drained soils retain a certain amount of moisture even through long periods of drought.

Other physical properties of soils that are of importance in soil corrosion are air-pore space, apparent specific gravity, water-holding capacity, moisture equivalent, and shrinkage. The air-pore space is the percentage of the volume of soil at a definite moisture content that is occupied by air. It is therefore a relative measure of the permeability of the soil to air and of the rate at which moisture can move through the soil. Other things being equal, a high value of air-pore space indicates a relatively noncorrosive soil. The apparent specific gravity, which is the weight of a unit volume (1 ml) of undisturbed soil, is an index of the compactness of mineral soils because the true specific gravity of the mineral particles in soils varies only within narrow limits. This

value is influenced to a large extent by the amount of voids in the soil. The moisture equivalent of a soil is defined as the percentage of water retained by a previously saturated layer of soil of given weight when subjected to a centrifugal force of 1,000 times gravity. The moisture equivalent is a measure of the retentiveness of soils for water. The quantity of water retained by soils when equilibrium with capillary and gravitational forces has been established is the water-holding capacity of the soil.

The volume shrinkage is a measure of the colloidal nature of the particles in a soil and indicates the tendency of the soil to crack on drying and to swell when wetted. The cracking of soil permits more oxygen to reach buried pipe and so affects the character of the corrosion products and the differential aeration potentials. Some soils on drying form hard clods which at times act much like stones with respect to differential aeration and pressure on pipe coatings. Some soils adhere tightly to certain pipe coatings, and as they shrink on drying tend to pull the coating from the pipe [41]. Some coatings are pressed into the soil cracks when the soil expands due to the absorption of water. Figure 3 shows cracks in Lake Charles clay soil and ridges in a rag-felt-reinforced asphalt coating caused by soil pressure. Thus soil shrinkage is important to corrosion and pipe-line protection. In general, clay soils and soils high in organic matter shrink the most, but not all clay soils shrink greatly on drying. For example, Bell clay has a clay content of 44.9 percent and a shrinkage value of 23 percent, whereas Susquehanna clay has a clay content of 45.8 percent and shrinks only 4.7 percent. Fargo clay loam has a clay content of 70.1 percent and a shrinkage of 21.0 percent.

The physical properties of soils are described by Keen [42]. This publication, however, does not discuss the relations of the physical properties of soils to corrosion. The chemical and physical properties of soils, as well as their origin and classification, are discussed in the Atlas of American Agriculture [33] and Soils and Men [43, 44], but neither of these books discusses soil corrosivity.

The physical characteristics of soils that are favorable to corrosion are poor aeration, and high values for moisture equivalent, clay content, apparent specific gravity, and shrinkage. These are characteristics of heavy, poorly drained soils. They are not independent of each other, and their effects on corrosion may be modified by the chemical characteristics. Table 5 shows the relative corrosivity of some of the soils in the National Bureau of Standards tests, as indicated by the average of four criteria [45] and the values of some of the physical properties of those soils.

Soils differ greatly in corrosiveness, and their corrosivity is affected by many soil properties. Table 6 shows the maximum and minimum values of certain data on soils at National Bureau of Standards test sites. The range of values for all soils in this country is considerably greater. Widely different soils frequently are found within a few hundred feet of each other, and corrosive soils are found in most parts of the United States.

As soil characteristics differ greatly and corrosive soils are widely distributed, it is obvious that no average value for the corrosiveness of soils and no allowance or factor of safety based on average soil conditions should be used in the design of a pipe line to be installed under specific soil conditions. It is necessary in each case to consider the cor-



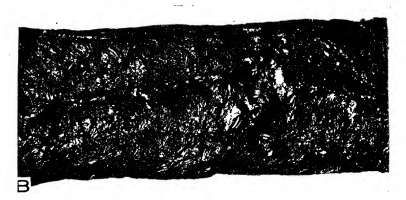


FIGURE 3 .-- Soil cracking and its effect on a pipe coating.

A, Cracks in Lake Charles clay. The white spot below the junction of the cracks is a fifty-cent coin; B, rag felt reinforced asphalt pipe coating wrinkled by soil prossure. Swelling of the soil forced the coating into the cracks. Note also cracks in the coating.

rosiveness of the soils that the pipe line will encounter if proper account of soil corrosion is to be taken. Any other procedure must result in waste on parts of the line and inadequate protection for other parts.

As many of the factors that influence corrosion are not closely related and do not vary together, one cannot expect to find a close correlation between corrosion and any single soil characteristic. It is important to remember this when soil surveys and corrosivity tests are considered.

# IV. INVESTIGATIONS BY THE NATIONAL BUREAU OF STANDARDS

### 1. GENERAL PLAN AND METHODS

In 1921 the American Committee on Electrolysis, of which the National Bureau of Standards was a member, decided that the effect of

Table 5.—Correlation of soil corrosivity with the physical properties of soils.

Relative corrosive- ness	Soil number	Moisture equivalent (compacted soil)	Aeration ^a	Air-pore space	Apparent specific gravity	Volume shrinkage	Resistivity at 60° F (15.6° C)	
1 2 3 4 5	^b 28 43 14 4 40	Percent 19.6 (c) 12.2 22.2 31.0	VP VP F P P	Percent 2.5 (c) 14 4 7.0 2.3	(c) (e) 1.76 1.78 1.78	Percent 5.9 (c) 1 0 2.2 16.4	Ohm-em 408 60 3,520 6,670 970	
6	42	24.8	F	14.9	1.79	4 7	13,700	
7	37	7 0	F	(e)	(e)	0	11,200	
8	8	34.8	P	8.7	1.56	21.0	350	
9	11	31 3	G	15.5	1.49	8.6	11,000	
10	16	16.5	F	12.0	1.65	0.6	8,290	
11	39	18.3	P	7.5	1.72	3.8	7,440	
12	41	28.1	F	6.9	1.61	14.6	1,320	
13	44	25.3	G	7.2	1.55	6.0	1,000	
14	3	29.9	G	18.2	1.60	7.0	30,000	
15	30	24.0	P	7.2	1.81	7.5	1,500	
16	32	11.8	G	11.7	1.85	0.1	5,700	
17	19	26.3	F	3.9	1.76	11.8	1,970	
18	2	35.2	P	2.0	1.95	23.0	684	
19	22	28 4	G	9.6	1.67	3.0	5,150	
20	18	22.0	G	16.6	1.26	1.3	1,410	
21	25	18.6	F	9.5	1.95	7.6	1,780	
22	7	36.4	P	3.7	2 02	34.5	2,120	
23	36	14.9	G	16 0	1 62	0	11,200	
24	17	27.7	P	4.4	1.72	5.4	5,980	

" VP = very poor; P = poor; F = fair; G = good.
b Most corrosive soil.
Not determined.

Table 6.-Maximum and minimum values of certain properties of the soils at the National Bureau of Standards test sites.2

Property	Maximum value	Minimum value	
Electrical resistivity Ohm-om  Moisture equivalent 76 Air-pore space 76 Apparent specific gravity 77 Volume shrinkage 76	54,400 75.5 40.6 2.08 42.7	32 2 3 1.1 1.41	
Total aciditymg-eq/100 g of soil. pH Annual precipitation ^b in. ² Mean air temperature ^b o°F Composition of water extract:	297.0 10.2 61.6 71.8	0 2.6 2.0 39.	
Na + K as Na	$\begin{array}{c} 45.1 \\ 19.24 \\ 9.45 \\ 4.6 \\ 2.1 \\ 43.3 \\ 46.5 \end{array}$	°0 0 0 0 0	

Determinations by I. A. Denison, R. B. Hobbs, and I. C. Frost.
 Data furnished by United States Weather Bureau.
 Zero values are estimated from the specific resistance of the soil.

soils on pipe unexposed to stray electric currents should be investigated, and the Bureau undertook the project. Soon thereafter the committee became inactive, and some years later it disbanded.

The original purpose of the soil-corrosion investigation was to determine the extent to which soils free from currents from street railways and other sources of power were corrosive with respect to the metals commonly placed in them. It is important to remember this objective, as it explains many things with respect to the conduct of the investigation. The reader should also bear in mind that many phenomena related to the behavior of metals in soils were not generally understood at the time the tests were undertaken.

The next step in the investigation was a meeting of the technical representatives of a number of manufacturers of cast and wrought ferrous pipe with representatives of the National Bureau of Standards. At this meeting an agreement was reached as to the sizes and varieties of materials for the initial tests and the way in which the specimens should be buried. The Bureau of Soils, now the Bureau of Plant Industry, Soils, and Agricultural Engineering, of the United States Department of Agriculture, acted as an adviser in the selection of the types of soils to which the specimens should be exposed and assisted in the selection of the test sites in order that, insofar as was practicable, the sites should represent definite soil types. However, in order to secure labor and keep down expenses, it was sometimes necessary to use sites that were not altogether satisfactory, and which might have been rejected had the importance of very uniform soil conditions been fully realized.

Each time specimens were removed, manufacturers and other cooperators were invited to have representatives present to note soil conditions and the condition of the specimens. On each occasion one and usually several cooperators' representatives participated in the removal of the specimens. A rough draft of the report on the specimens was submitted to each manufacturer concerned, and usually a conference was held for the purpose of discussing and revising the report. In this way the Bureau obtained the advice and suggestions of a large number of experts in different phases of corrosion. With a very few exceptions, the revised reports had the approval of all concerned. The so-called National Bureau of Standards soil-corrosion investigation is therefore more than the name implies, and the results can be accepted with a considerable degree of confidence, provided the way in which they were

obtained is fully understood.

The cooperators in the investigation are listed in Appendix 1. Each cooperator has been listed only once, although many of them have cooperated in several ways and in several tests. The soils where the

specimens were buried are described in Appendix 2.

Figures 4 to 6 illustrate the ways in which the specimens were placed in the trenches. The depths at which the specimens were placed corresponded roughly to the depths at which pipes had been laid in the same region, and ranged from 18 inches in the southern locations to 6 feet in northern locations. Before the specimens were buried, careful tests were made for stray electric currents. The specimens were placed in a definite order with respect to each other and from 6 inches to 1 foot apart to avoid one specimen affecting the corrosion of another and to facilitate the removal of one set of specimens without disturbing the soil near the other sets. When the specimens were removed, the upper side or edge of each was marked with a center punch. This made it possible to study the distribution of corrosion on each specimen with respect to its position in the trench, and with respect to other specimens.

As a rule, 10 or 12 specimens of each material were buried at each test site, and 2 specimens were removed at intervals of approximately 2 years. In later years the times between removals were modified in accordance with the apparent corrosiveness of the soil to which they were exposed. The removal of specimens after different periods of exposure was to obtain data on changes in rates of corrosion. It was hoped that the life of a pipe of a given wall thickness could be predicted from these

changes.



FIGURE 4.—Burial of soil corrosion specimens near Alexandria, Va., in 1932.

After the specimens were removed from the trench, the loose dirt was scraped off and the specimens were boxed and returned to the National Bureau of Standards. Most specimens having protective coatings terminated in caps that supported the specimens and prevented the coatings from touching the box or each other during shipment. When the specimens reached the laboratory, each different material was subjected to appropriate chemical and mechanical treatments, to remove the corrosion products with an insignificant loss in weight of the base metal and without mechanical injury to the specimen. The methods employed in cleaning the specimens are described in detail in Appendix 3, and the methods of obtaining, computing, and reporting the data are described in Appendix 4. Careful tests showed that the wrought ferrous specimens lost very little metal by the cleaning processes. The penetration of corrosion on cast iron is along the boundaries between the graphite and the metal crystals and some uncorroded iron probably was removed with the corrosion products. On the other hand, tests showed that usually not quite all the corrosion products were removed by cleaning the specimens.

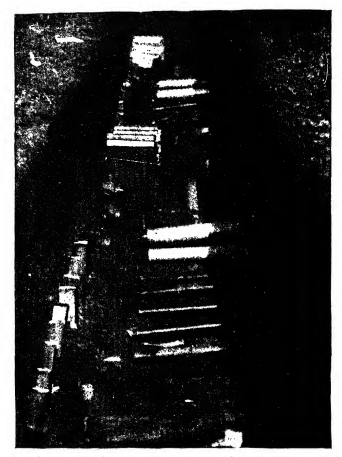


FIGURE 5.—Arrangement of specimens buried in 1934
Note two soil horizons.

The soil-corrosion investigation started with the burial of only the most commonly used pipe materials and coatings. From time to time, as specimens were removed, other materials were added to the tests. By 1932 it had become evident that the corrosion of the commonly used ferrous pipe materials was rapid in some soils, and a new series of tests was started for the purpose of finding materials suitable for use in corrosive soils. For these tests 15 corrosive soil test sites were chosen, and manufacturers were invited to submit materials which they believed suitable for corrosive-soil conditions. This considerably increased the already long list of materials under observation and introduced a few materials that were unsuitable for testing by the methods applied to ferrous pipe. New pipe coatings were also submitted, although previous tests had shown that tests of coatings applied to short lengths of pipe served only



FIGURE 6.—Arrangement of specimens buried near Baltimore, Md., in 1932.

to eliminate definitely unsuitable coatings and were insufficient to identify coatings that would effectively protect operating pipe lines.

### 2. MATERIALS TESTED

Tables 7 to 18 show the form, dimensions, composition, and the number of specimens of each material submitted for the National Bureau of Standards tests, and also the coatings tested by the American Gas Association (AGA) and the American Petroleum Institute (API) under the direction of the Bureau. Tests have been started on more than 36,000 specimens. At present, about 6,500 specimens remain exposed to 17 soils.

It will be noted in the tables that the same letter has been used to designate different kinds of materials. The identification of the material involves an identifying letter, the name of the class of materials to which the specimen belongs, and the year in which the specimen was buried. The form, dimensions, and age of a specimen are also helpful for determining the group to which the specimen belongs. As the specimens are grouped as to age and material in all the soil-corrosion reports, readers of the reports should have little difficulty on account of the identification of the specimens.

	I		T		
- Material	Iden- tifi- cation	Year buried	Number of speci- mens buried	Form	Nominal width or dia- meter
Wrought iron: Hand-puddled. Do Do Do Co Roe process. Carbon steels, wrought iron and pure open-bearthiron:	b, d B, D A B	1922 1922-28 1932 1932	564 742 150 150	Pipe do do do	in. 1.5 3 1.5
Pure open-hearth iron. Open-hearth iron Do Open-hearth steel. Bessemer steel. Do Low-carbon steel.	A A A K e Y M, N	1924 1922 1928 1922 1922 1922 1922-28 1932	40 568 174 568 568 568 588 150	Sheet Pipe do do do do do do	2.0 1.5 3 1.5 1.5 2.5
Copper-bearing steel  Special open-hearth steel Copper-molybdenum open-hearth iron	S A M, M	1922 1924 1937	568 40 150	Pipe Sheet Plate do	3 2.0 2.5 2.5
Do Do Copper-nickel steel Nickel-copper steel Do Do	OXHHI BYD	1937 1937 1932 1941 1937 1937 1941 1932	150 150 150 150 150 150 150 150	do do Pipe do Plate do Pipe do	2.5 2.5 1.5 1.5 2.5 2.5 2.5
Chromium-silicon-copper phos- phorus steel 2% chromium steel with molyb-	C N. N	1937 1939	150	Plate	2.5
denum. 2% chromium steel with molyb- denum.	K, K	1939	150	Pipe Plate	1.5 2.5

4 to 6% chromium atcel	2.3 2.5 2.5
4 to 6% chromium steel with 15 1937 150 Plate 1937 150 do	-
molybdenum E 1937 150 do	2.5
4 to 6% chromium steel with	2.5
4 to 8% chromium steel with	- 1
	2.5
	4 1.5
26% chromium C 1932 60 Pipe	* = 1
	1.0
	- 1
18% chromium steel with nickel K 1932 150 Sheet	3
	1.5
18% chromium steel with nickel W 1932 175 Sheet	4.0
and insurance with nickel	- 1
	6
	6
18% curomium steel with nickel.	- }
manganese and molybdenum. C, M 1939 150 Plate	2
	1
	2.5
and manganese Y 1932 175 Sheet	1

re ar

he Na na

Sla

Length	Thick- ness	С	Si	Mn	s	P	Cr	Ni	Cu	Мо	Other elements
in.	in.	%	%	%	%	%	%	%	%	%	%
6 6 12 12	$\substack{0.145 \\ .216 \\ .145 \\ .145}$		.15	Trace 0.033 0.29 0.41	$0.023 \\ .022 \\ .018 \\ .018$	. 195 . 160			.03		Oxide +slag 2.56
6 6 6 6 6 6	.125 .145 .216 .216 .145 .145 .216 .145	.02 .02 .12 .09 .08 .15		Trace do .41 .39 .38 .40 .49	.05 .05 .036 .046 .050 .038	.088			.014		F ₀ 99.9
6 6 12	.216 .062 .188	.07		. 24 	.032 	1	.1	o. o.			O2, 0.015
12	.25	.04		. 16	.027	.008	.04	. 14	.05	10.0	7 N2, 0.008 Sn, 0.002
12 12 12 12 12 12 12 8.7	.25 .25 .145 .145 .265 .248 .154 .145	.03 .06 .04 .03 .06 .07 .12 .14	.047	.098 .32 5 .16	.03: .02: .02: .02: .02: .02:	0 .06 7 .01 5 .00 5 .09 2 .01	9 .02	. 52 1.96	54 .52 .47 .95 1.01 1.04	.0	7
12	. 188	.07	5 .84	.20	.01	s .12	4 1.02	0.22	0.42	s¦	
14.8	. 145	.09	.25	46	.01	0 .01	<b>5</b> [1.93	<u>'</u>			9
12	.175	l .os	2 .51	.46	.01	5.01	7 2.01	.07	1.00	41.5	7

$^{10}_{12}$	. 151 . 245	:13		.46	.025 .005	:012 :013	5.05 5.02	.09	: ioos	. : : .	
12	. 188	.074	.41	.32	.006	.013	4.67	.09	.004	.51	Al. 0.030 Ti, 0.022
12	. 203	.060	. 39	.40	.014	.021	5.76	.17	.004	.43	Al, 0.27
6 12 6	.063 .063 .145 .250	.065 .070 .12	.28 .34 .277	.38 .36 .42	.017 .015 .017	.014	11.95 17.08 17.72 26	.482 .092 .287	.021		
$\substack{\substack{11 \\ 12 \\ 6}}$	.025 .145 .063	.08 .05 .093	.33 .28 .42	.44 .46 .36	.022 .011 .017	.015	17.20 17.52 18.69	8.95 8.85 9.18	 	::::	
10	. 068	.06	.40	6.09			17.76	3.83	.95		1
10	. 063	.07	.48	9.44			17.78		.74		
12.5	. 25	.07	.40	1.24	.008	.016	17.78	10.96		2.63	
12	. 25	.07	.91	1.99	.012	.014	19.27	22.12	1 07	3.52	
6	.063	.144	. 59	1.80	.011	.015	22.68	12.94	.021		

rg Corrosion

ಲಾ

TABLE 10.—Composition of sino, lead, and aluminum-aloy materials.

						-	-	-	-	-	-	-		-	_	_		
Material	Identi- fication	Year	Number of speci- mens in test	Form	Nom- inal width or di-	Nom- inal width or di- ameter	Thick- ness	Zn	T¥	C _u	Fe	Pb	. Sh	Bi	<del>2</del> 8	:55	Mn	Other elements
					iñ.	ii.	in.	%	%	%	1%	%	%	2%	%	%	5%	200
מני: סיווים הייה	Z	1937	150	Plate	2.3	12	0.15	<u> </u>	÷	0.009 0.095	600	0.095	:	-		<del>-</del> -	:	0.0038 Cd. (0.003 Cd.
Die casting zine	Z	1937	150	ę	4.44	8.9	.122	:	4.00 1.05	.05	.018	× .003	.018 < .003 < 0.001	:	<del>-</del> -	-	-	0.02 to (0.05 Mg.
Standard zinca. Zinc. Do.	$_{2}^{\mathrm{P}}$	1924 1924 1924	40 40 40	Sheet do Plate	2.0 2.0 2.0	999	.062	+ 2.5. 89.5. 99.5.										
Hearth refined lead Hearth refined lead Chemical lead Do Tellurum lead Artimonial lead Commercial lead	HEALTONN	1924 1924 1937 1937 1922 1937 1922	40 40 150 150 281 281 281	Plate do Pipe do Sheet Pipe Sheet	8.5.5.00 3.5.5.5.00	222222 2222222	.25 .25 .177 .177 .112			0.002 .57 .056 .082 .088 .036		99.99 99.93 99.94 99.87 99.12 99.64		0.004 .002 	0.004 .002 0.0011 .0016 5.31			0.009 Ag. 043 Te.
Lead alloy	M	1941	150	Pipe	1.25	15	.172		:				0.25				<u>:</u>	plus usual impurities in lead.
Aluminum: Comnercial aluminum Aluminum with manganese Duralumin	చెచిచి	1924 1924 1924	0 <del>4</del> 40	Sheet do do	2.0 2.0 2.0	000	.062		0.02 97.75 0.02 97.75	.09 1.16 1.10	. 30 . 30					0.3 #. 23	0.03 1.12 0.62	0.52 Mg.

a Standard composition.

Table 11 .- Metallic-coated specimens.

Material	Identi- fica- tion	Year buried	Number of speci- mens buried	Form	Nominal width or diameter	Length	Wall thick- ness	weight of	Average thick- ness of coating
Zinc coatings:  Bessemer steel Pure open-hearth iron Wrought-iron Do Copper-bearing open-hearth	B A D D	1922 1923 1923 1923	124 208 24 28	Pipe do do do	in. 2.0 2.0 3.0 1.5	in. 24 17 17 17	in. 0.154 .154 .216 .145	$0z/ft^2$ 1.81 2.82 3.48 4.99	in. 0.0032 .0050 .0061 .0088
copper-bearing open-nearth steel.  Pure open-hearth iron.  Do.  Copper-bearing steel.  Do.  Do.  Do.  Bessemer steel  Pure open-hearth iron.  Bessemer steel.  Copper-bearing steel.  Steel.  Lead coatings:	Y A2 A3 A4 Y2 Y3 Y4 Y4 CA bCA bCB cT	1923 1924 1924 1924 1924 1924 1924 1924 1924	56 28 188 28 28 188 28 188 56 56 56 300	do Sheet do Pipe	3.0 66 66 66 66 66 66 67 57	17 12 12 12 12 12 12 12 12 12 12 12 14	.216 .062 .062 .062 .062 .062 .062 .062 .050 .050	3.47 1.79 1.98 2.65 1.57 2.15 2.76 2.92 1.62 1.87 1.66 2.12 3.08	.0061 .0032 .0035 .0047 .0028 .0038 .0048 .0051 .0028 .0033 .0029 .0037
Steel	L CA	1924 1932	376 150	do do	1 5 1.5	6 13	. 145 . 145	:::::::	.0025 .0015
Aluminum coatings: Wet calorized Dry calorized	::::::	1924 1924	48 56	Pipe do	2 0 2.0	6	. 154 . 154		::::::
Tin coating: Tinned copper		1937	150	do	1.5	12	.060		

<sup>a 1 oz/ft² of surface =0.00172 in.
b Uncoated black sheets buried for controls.
e Black iron pipe buried for controls.</sup> 

 ${\tt Table \ 12.--Nonbituminous-nonmetallic \ coatings.}$ 

Identi- fica- tion	Year buried	Number of speci- mens		${f Description^a}$
•	1932	150		A synthetic rubber, stated to be an olefin polysulfide reaction product, was exposed in the form of sheets 10 by 5 by 11 inch. Subsequently, a process was developed which permitted application of this material to pipes. Specimens coated by this process were exposed in 1939.
C	1932	150	0.014	Two coats of porcelain enamel, one of which was acid-resisting.
D	1932	150	.010	First coat, 23-percent solution of a rubber derivative in xylene; second and third coats, 30-percent solution of the rubber derivative in xylene; fourth coat, 20-percent solution of the rubber derivative in a mixture of turpentine and mineral spirits. Five percent of the solids was carbon black.
E	1932	150	.005	Two applications of paint that differed in color. Neither the kind of pigment nor the kind of vehicle was specified.
F	1932	150	.060	A semiplastic compound, which was applied cold with a brush, consisting of 4½ parts of treated cashew-nutshell oil, 3 parts of asbestos fiber, and 3½ parts of mineral turpentine substitute.
G	1932	60	.090	A hard-rubber compound, containing rubber, sulfur, and an accelerator, cured to a bone-hard condition.
н	1932	60	.100	A highly loaded hard-rubber stock which contained 30 percent of magnesium carbonate and approximately 15 percent of "white substitute."
J	1932	30		A synthetic resin varnish baked at 425° F for 30 minutes.
K	1932	30	.006	A paint containing highly chlorinated rubber, dissolved in a suitable solvent, to which may have been added drying oils, pigments, quarts meal, or carborundum. This coating was applied in Germany to pipe 1 inch in outside diameter.
w	1932	150	.170	An experimental coating prepared as follows: The pipe was primed with a china-wood-oil varnish containing zinc chronate and basic lead chromate and basic 200° F for ½ hour. A top coat of dehydrated china-wood oil containing powdered mica and a catalyst was molded on the pipe and heated to 200° F for 3 hours.
L	1937	150	.007	Two coats of a Bakelite varnish followed by two coats of another type of Bakelite varnish, each coat being baked after spraying. It appears as though the baking has resulted in a fusion of the separate layers.
$M_1$ .	1937	150	.004	Two coats of a Bakelite priming followed by two coats of a bakelite chemically resistant aluminum paint. Facth coat was sprayed and allowed to air dry overnight. More specifically the priming consisted of a Bakelite 25-gallon varnish—55 parts, zinc chromate, 30 parts, asbestine, 15 parts, to which was added lead-cobalt-manganese drier. The finishing aluminum paint consisted of 2 pounds of aluminum paste to a gallon of a Bakelite chemically resistant varnish.
M2.	1937	150	.003	The same as coating M ₁ , except that it was applied to 2½ by 12-inch plates instead of to pipe.
N	1937	150	.032	A double layer of Bakelite-treated asbestos tape, the second layer overlapping the fir t, which was made to adhere to the pipe and to the first layer of tape by the use of an anticorrosive resin compound. A final protective coat was used over this tape, consisting of a spray coat of the same Bakelite chemically resistant aluminum enamel as used on the Mispecimens.
	1937	50	.033	A fabric coated on one side with Koroseal to an over-all thickness of 0.03 inch. The fabric was wrapped spirally on the pipe under tension with the Koroseal next to the metal and then painted with two coats of Koroseal solution applied hot.
R	1939	150	.0055	A blue basic lead sulfate phenolic resin varnish paint consisting of two coats of the following composition:  Basic lead sulfate in raw linseed oil (90% pigment)
s	1939	150	.044	Plastic made of pure vegetable gum base with nondrying oils and asbestos fiber applied cold. Shielded by a spiral wrap of impregnated cotton fabric.
т.	1939	45	.035	Thiokol sprayed on.

^a The coatings were applied to 1½-inch standard pipe, 12 inches in length, unless otherwise stated. The descriptions of the coatings were furnished by the manufacturers.

Table 13 .- Miscellaneous specimens included in the National Bureau of Standards tests.

Symbol	Material	Year buried	Number of specimens buried
A B C D	Decarburized, malleable-iron nuts and bolts. Not decarburized, malleable-iron nuts and bolts. High-strength, malleable-iron nuts and bolts Steel nuts and bolts. Charcoal east-iron nuts and bolts.	1932 1932 1932 1932 1939	48 48 48 48 150
CE E	Steel nuts and bolts. Sheradized nuts and bolts. Lead-coated nuts and bolts. Black wrought-iron nuts and bolts. 2-inch cast steel elbows.	1939 1924 1924 1924 1924	150 100 96 100 56
I S P	4-inch machined cast-iron pipe 2-inch semisteel nipples. 2-inch malleable-iron bends. 1½-inch couplings attached to threaded pipe. Composite steel plates.	1924 1924 1924 1922 1941	24 48 48 192 150
С С	Miscellaneous Bureau of Mines zinc plates 1-inch chrome iron pipe 2-inch copper-steel pipe 2-inch machined wrought-iron pipe 2-inch unfinished wrought-iron pipe.	1924 1926 1926 1926 1926	27 66 20 18 18
L M P	2-inch leadized pipe.  2-inch machined Bessemer steel pipe.  2-inch unfinished Bessemer steel pipe.  7- Four different mixtures of pipe fittings (brass caps attached to lead, brass, and galvanized nipples).  11 types of bituminous coatings.	1926 1926 1926 1924 1924	42 18 18 2,208
	Armoured cable American Gas Association bituminous-coating tests* American Petroleum Institute bituminous-coating line tests* American Petroleum Institute bituminous-coating nipple tests* Asbestos-cement flue pipe	1924 1929 1930 1930 1932	246 2,352 2,050 2,208 150
·i	6-inch asbestos-cement pipe, class 150. 4-inch asbestos-cement pipe, class 150. Cement-coated east fron.	1937 1939 1924	150 150 52

^{*} See tables 17 and 18 for description and characteristics of these coatings.

b See tables 14, 15, and 16 for description and characteristics of these coatings.

Table 14.—Coatings in American Petroleum Institute line tests.

		and designation of the second
Symbol	Thickness	Description
A	in. 0.0647	Two coats of asphalt emulsion.
в	.1072	One coat of grease, spiral wrap of grease-saturated fabric (Osnaburg type), and outer coat of heavier-consistency grease.
C	.0206	Two coats of filled cutback coal tar.
E	.1506	Asphalt primer, followed by sling coat of asphalt compound, spiral wrap of coal-tar-saturated Osnaburg fabric, outer coat of asphalt and kraft paper.
F	.4185	Two coats of asphalt emulsion, followed by rigid shield of sand and cement mortar.
G	.0625	Coal-tar primer, followed by coal-tar -asphalt enamel, and unbonded wrap of asbestos pipe-line felt.
н	.0807	Coal-tar primer, followed by coal-tar enamel, and unbounded wrap of $w_{00\mathrm{d}}$ veneer.
ĸ	.0685	Coal-tar primer, followed by coal-tar-asphalt enamel.
L	.0798	Coal-tar primer, followed by coal-tar enumel.
м	.0576	Do.
N	.0596	Coal-tar primer, followed by coal-tar-asphalt enamel.
0	.5186	Asphalt primer, followed by a hot coat of priming asphalt, and spiral wrap of asphalt mastic carried on pipe with tissue and sheathing paper (machine-applied).
R	.1427	Asphalt primer, followed by two coats of asphalt enamel; spiral application of asbestos pipe-line felt; flood coat of enamel, and kraft paper (machine applied.)
s	.1502	Asphalt primer, followed by two coats of asphalt; spiral application of rag-base pipe-line felt; flood coat of asphalt, and kraft paper (machine-applied).
т	.3507	Coal-tar primer, followed by two straight-away rag-base pipe-line felt applica- tions on the inner faces of which coal-tar enamel mopped on; sling coat of coal-tar enamel, and whitewash.
U	.1709	Coal-tar primer, followed by two coats of coal-tar-asphalt enamel; spiral application of asbestos pipe-line felt; flood coat of enamel, and kraft paper (machine-applied).
x	.2302	Hot asphalt primer, followed by double-spiral wrap of unsaturated fabric (Osnaburg type) drawn through molten asphalt, and spiral-butt wrap of 26-gage strip steel.
Y	.0287	Asphalt primer, followed by one coat of asphalt cutback; one coat of asphalt adhesive, and machine-wrap of aluminum foil.
z	. 2062	Hot asphalt primer, followed by double-spiral wrap of unsaturated fabric (Osnaburg type) drawn through molten asphalt, and kraft paper (machine-applied).

Table 15.—Coatings on American Petroleum Institute short pipe sections.

Symbol	Sponsor	Average thickness	Description
a	American Tar Products Co	in. 0.054	Pyrmax primer, followed by one coat of Komac P. C. enamel (coal tar).
b	Arco Co	.020	Two coats of filled Arco Q. D. Savaline (cutback coal tar).
bb	do	034	One coat of filled Arco Q. D. Savaline; spiral wrap of saturated fabric (Osnaburg type); and final coat of Savaline.
c	Barber Asphalt Co	.072	Asphalt primer, followed by two spiral wraps of saturated fabric (Osnaburg type), and kraft paper—Genasco pipe-coating type 1—(fabric and kraft paper machine-applied).
ce	do	.045	Two coats of cutback asphalt—Genasco pipe- coating type 2.
d	The Barrett Co	.342	Barrett coal-tar primer, followed by two coats of rag-base rooling felt to the inside face of which hot Barrett pipe-line enamel (coal tar) was mopped on, and an outer coat of enamel fol- lowed by whitewash.
4d	do	. 050	Barrett coal-tar primer, followed by Barrett pipe- line enamel (coal tar) and whitewash.
ddd	do	a.050	Barrett coal-tar primer, followed by Barrett pipe- line enamel (coal tar) and unbonded polymer- ized resin shield.
e	Dearborn Chemical Co	.026	One coat of No-Ox-Id G special, spiral wrap of grease-saturated fabric (Osnaburg type), and outer coat of service coat (heavier-consistency grease).
ee	do		Same as e excepting impregnated ashertos-base roofing felt used in place of woven tabric.
f	E. I. duPont de Nemours and Pire Nove (1)	.149	Two coats of asphalt-chromate emulsion, spiral- butt wrap of asbestos pipe-line felt, outer coat of A.C.E. and whitewash.
ff	do	.190	Two coats of asphalt-chromate emulsion followed by a coat of sand and cement mortar.
fff	do	.037	Two coats of asphalt-chromate emulsion (white-washed).
g	Eagle Picher Lead Co	.069	Under coat of sublimed blue lead followed by bitumastic primer (coal tar) and bitumastic XXH (coal-tar enamel). See coating zzz.
h	Benjamin Foster Co	069	Asphalt primer followed by one sling coat of I.B.M. bituminous pipe coating No. 4 (asphalt mastic).
j	Headley Emulsified Products	.015	
k	CoIIill, Hubbell and Co	.075	Three coats of Headley No. 11 asphalt emulsion. Biturine primer, followed by sling coat of biturine No. 212 enamel (coal-tar-asphalt).
kk	do	.165	Biturine primer followed by two coats of biturine No. 212 enamel (coal-tar-asphalt), spiral wrap of J.M. 15-lb asbestos pipe-line felt, flood coat of enamel, and kraft paper (machine-applied).
1	Inertol Co	.031	Inertol No. 49 primer followed by two coats of seal coat (cutback asphalt) and kraft paper (kraft paper machine-applied).
m	Andrew McLean Co		Priming coat of zinc chromate, followed by McLean's asphalt emulsion and wrapped with asphalt-saturated burlap and kraft paper (burlap and kraft machine-applied).
n	Everlast, Inc	.158	McEverlast penetration followed by a coat of hot blended compound (asphalt), spiral wrap of coal-tar-saturated Osnaburg fabric, outer coat of H.B.C. and kraft paper (fabric and kraft paper machine-applied).
nn	do	.337	Asphalt primer, followed by hot coat of priming raid at a spiral wrap of Somastic (asphalt rasic, carried on pipe with tissue and sheathing paper (machine-applied).
p	Lead Industries Assn	.071	One coat of red-lead paint, followed by bitumastic primer and bitumastic XXH (coal-tar enamel). See coating zzz.
pp	do	.138 .015	One coat of red-lead paint followed by coating q. Priming coat of red-lead paint followed by two coats of Headley No. 11 (asphalt emulsion).

Table 15.—Coatings on American Petroleum Institute short pipe sections —Continued.

Symbol	Sponsor	Average thickness	Description
q	National Tube Co	in. .155	Hot Robertson asphalt, followed by double-spiral wrap of unsaturated fabric (Osnaburg type) drawn through molten Robertson asphalt and kraft paper (machine-applied).
qq	do	216	Same as coating q, excepting 26-gage strip steel substituted for kraft paper.
qqq	do	.183	Hot Robertson asphalt, followed by extruded National mastic (asphalt).
r	Fish-Schurman Corp	. 105	One coat of grease, followed by spiral-wrap of Denso-Tek protective tape (grease-saturated Osnaburg-type fabric), and outer coat of asphalt-saturated burlap.
s	Resistcor Engineering Corp	.041	Resisteor primer (coal tar) followed by sling coat of Resisteor enamel (coal-tar-asphalt).
ss	do	a.041	Resisteor primer (co.d-tarsp\ all) followed by sling coat of Resisteor crarted conditar) and unbonded wrape of M. 15-lb usbests pipe-line felt.
t	Sherwin-Williams Co	.052	Lynkote primer (coal tar), followed by sling coat of Lynkote enamel (coal tar).
u	James B. Sipe and Co	.047	Impervobond No. 411-083 (asphalt primer), followed by cutback-asphalt compound sprayed on.
v	Standard Oil Co. of California.	.337	Same as nn, using asphalt-mastic pipe coating.
vv	do	. 079	Asphalt primer, followed by Sonamel (asphaltenamel) tissue paper, and kraft.
w	Standard Oil Co. of New Jersey.	. 092	Asphalt primer and two coats of Standard pipe coating (asphalt-enamel).
ww	Johns-Manville Corp. and Standard Oil Co. of New Jersey	.170	Asphalt primer, followed by one coat of Standard pipe coating (asphalt-enamel), spiral application of J.M. 15-lb asbestos pipe-line felt; flood coat of enamel, and kraft paper (fabric machine-applied)
x	Technical Products Co., Inc., Aluminum Co. of America	.018	Tec-Pro primer (asphalt), followed by one coat of Tec-Pro black line No. 45 (asphalt cutback) and Tec-Pro adhesive (asphalt) and machine- wrap of aluminum foil.
у	The Texas Co	.144	Texaco primer (asphalt), followed by two coats of Texaco No. 30 asphalt pipe coating, spiral wrap of Texaco pipe-line icht (asphalt-sat urated rag- base); flood coat of asphalt and kraft paper (machine-applied).
уу	do	.199	Texaco primer (asphalt), followed by two coats of Texaco No. 30 asphalt pipe coating, spiral wrap of Texaco pipe-line felt; coat of asphalt, second ply of felt and kraft paper (machine-applied).
ууу	do	.259	Same as conting yy, excepting a flood coat of Texaco No. 30: Spin I: pipe conting over second ply of fe'r. Once wrap of Scatt paper.
уж	Udylite Process Co	.0004	Electrodeposited cadmium-udylite. Cadmium.
z	Wailes Dove-Hermiston Corp.	.072	Bitumastic primer, followed by sling coat of bitumastic XH enamel (coal tar) and white- wash.
zz	Wailes Dove-Hermiston Corp. Merchants Basket and Box		
	Co	*.077	Bitumastic primer (coal tar) followed by sling coat of bitumastic XXH enamel (coal tar) and unbonded wrap of Becker's sewed wood veneer.
zzz	-	.077	Bitumastic primer, followed by sling coat of bitumastic XXH enamel and whitewash.
zx	do	.012	Two coats of bitumastic No. 4 (eutback coal tar).

a Without shield, felt, or wood veneer.

Table 16.—Characteristics of bituminous coating materials in the American Petroleum Institute tests.

[Determinations by H. S. Christopher]

Designation of material	Specific gravity	Ring-and- ball softening point	Penetration (Dow) at 115° F in 5 sec (50 grams)	Ductility (Dow) at 115° F	Consistom- eter hardness (Abraham) at 115° F	Insoluble in CS2ª	Asha
E G, N. H, L. K, U.	1 028 1.420 1.628 1 266 1.418	° F 186 202 205 192 145	20 3 2 16 83	cm 14.0 0 7 3.9 16.0 60.0	14 7 67.1 54.8 23.5 44.5	Percent 2 12 42.9 55.4 29.1 37.4	Percent 0.98 26 5 37.8 18 9 22.9
S T X, Z a d, dd	0 996 1.350 0.997 1.300 1.360	194 191 238 241 190	16 	3.2 4.1 2.5 2.7 3.6	10.9 63.5 17 1 34.5 56.5	1.0 34.1 0.7 42.8 32.7	0.4 14.4 0.13 10.9 13.2
g, zz, zzz. h k, kk n q, qq	1.610 1.685 1.259 1.032 1.006	205 189 192 190 232	2 21 27 14 33	3.8 10.3 62.0 12.6 2.5	61.3 13.0 9.1 16.9 16.0	55.6 61.6 21.9 4.6 0.8	35.3 61.3 5.6 2.4 0.4
s, ss	1.421 1.420 1.050 1.470 1.011 1.638	195 158 320 + 163 189 222	2 25 30 	2.8 58.2 0.0 7.2 3.2 1.7	63.0 10 0 36 7 8 8 10.9 65.4	40 6 39.3 17.0 42.2 1.1 57.6	26.9 21.4 13.4 39.9 0.5 40 2

 $^{^{\}rm a}$  When differences between the insoluble in CS₂ and ash are greater than 4 percent, the difference is apparently due to free carbon (a constituent of coal tar).

Table 17. Descriptions of pipe contings in

Machine
Sling

Brush Machino

Sym-	Manufacturer or agent	Nature of base primer	Method of applying prinor	Trade name of primer	Method of npptying conting
А	Am. Tar Products	ъ	Brush	A, T, P, primer	Sling
BB:::	Barrett Co. Benjamin Foster	P P	Dipped, no other	B. P. L. primer	ii do iiand
CC:	Co. do	a	freatment. Brush-hot	I. B. M. No. 1	Machine
$_{\mathbf{E}}^{\mathbf{D}}$ $_{\mathbf{E}}^{\mathbf{D}}$ $_{\mathbf{E}}^{\mathbf{D}}$ $_{\mathbf{E}}^{\mathbf{D}}$	DearbornChem.Co. do Emulsion Process	g B	Brush dodo	No-Ox-Id A special No-Ox-Id G special E. P. No. 1	Hand do Brush
F	Consolidated Gas Co. of N. Y. Flintkote Co. and E. I. duPont de Nemours & Co.	a	Dipped and dusted with coment. Dipped vertically, no other treat- ment.		
FF	Headley Good Roads Co.	a a	Brush do	A. C. E. +cement Headley No. 11	Brush do
<del>J</del>	Hill, Hubbell & Co. Johns-Manville Corp.	p p	do	Inhiburine J. M. tar primer	Sling Machine
JJ	do	n.	do	J.M. asphalt primer	do
EL	Arco Company Barber Asphalt Co.	p a	do	Arco ravenite Plastic pipe etg. Genasco primer	Brush do Machine
MM:	McEverlast, Inc.	a. s.	do	Penetrationdo	Brush Sling and hand wrap
N.L NB NT	National Tube Co. Lead Indus. Asan. National Tube Co. . do. . Paraffine Co., Inc.	a a a	Dipped Brush Dipped do, Brush	Robertson No. 228 Red lead Robertson No. 228 . do	Machinedo Machine Extruded Pabco Mach.

Q	Albrecht Pagen-	l g	Wrapped, no other	1
R	stecher. Resistor Eng, Corp. Sherwin-Williams Co.	B	Brushdo	Resist, primer Lyne-Kote primer
48 T	do	R	:.do	Texaco primer
υ	Hill, Hubbell &	1>	do	Inhiburine
¥v∷	Iroquois Gas Corp.	a a	do	Robertson No. 274
w	do	B	do	Inertol Bitumastic solu- tion
WF WL WR	Hermiston Corp.	<u>p</u>	do	Red lead Bitumastic Solu- tion.
ww	Milwaukee Gas Light Co.	P	do	Pabco primer
ž:	Enst Obio Gns Co. Hill, Hubbell & Co.	n. P	No primer Brush	inhiburine

Final application kraft paper. b a, asphalt; p, pitch; g, grease.

American Gas Association tests.

	· ·				
Nature of buses cont- ingb	Trade name of conting	reinforcement if any	Nan hours to apply coating	Average tempera- ture of ap- plication	Aver- ngo Dick- ness
P P B B B B B	A. T. P. pipe cont- place Notational Pipel and fell I.B. M. pipe cont- I. B. M. No. 2 No-Ox-Id GN 4-Q Parafait	15-1), rag felt * 15-1) asbestes No-Ox-Id muslin	2 8 4.0 4.5 3.0 7.5 7.5 10.5 4.0	350 350 370 375 375 Cold 280 Cold	0.070 .038 .134 .014 .120 .054 .071 .047
a	A. C. E.		4.0	do	.082
a a p	A. C. E. + cement Headley No. 11 Biturine J. M. pitch No. 2	14-lb asbestos	4.0 3.75 2.0 6.7	do 470 350	.029 .016 .034 .163
a.	J. M. asphalt	do	7.5	475	.126
a a	Save-a-line P. P. C.	Saturated cotton sheeting.a	2.7	Cold do	.056
a	Electrolysis proof	Osnaburg	4.3	do	. 010
a a a a	National coating do do Robertson No. 230	Muslin and steel	6.0 6.25 6.0	380 380 382 340	.146 .117 .147 .131
a	Floatine	Pabco felta	1 2.2	350	.123

AC.	Schade's wrapper	Schade's wrapper	6.0	Cold	.051
1>	Resistcor enamel Lyne-Kote hot		2.0	400 420	.081
n n	Lyne-Kote cold Texaco No. 30	i5-lb rag felt"	3 0 8.3	Cold 450	.007 .150
1>	Biturine spec.	30-lb rag felta	8.6	425	.214
a	Robertson asphalt Std. N. J. brick filler.	Cheeseclothdo	2.6 8.0	420 410	:079
D	Bitumastic (XXH)	do	3:8	440 456	.041
p p	Bitumastic do	15-lb asbestos	6 7 8 2 3 . 5	425 450 420	.154 .052 .0505
P	ifloatine	Wood veneer Pabeo rag felt	3:3	420	:100
a P	Biturine spec.	Sheeting 15-lb asbestos*	0.3	372 430	:115

TABLE 18. Characteristics of American Gas Association coating materials.

		Por not	35.1	19.5			17.6		1.5	: :	17.6	: : :	16.4	: :
	Insol-	11.C.S.	36.4	24.x 0.65.4	0.T	0 2	333 7	1.1	4. 2.	62.0	39.9	0 1	52.	
	Ash		%1.8 19.0	0 4 X	8.9		16 2 14 5		ÿ -4;-	63.7	25.7	5.6.7	35.8	e n
	B, & R.	point	282 167	25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55 25.55	229 102 to 122	110 to 115 145 to 155	187 159	135	154	255 144	201	176 176 208	181	218
	l at	37º J	kg 5.4 16.0	-10 C	10.5	5.9	19.0		13.2	18.6	13.0	8.6	19.0	17.7
	Тепяйе м(гепдсы	₹ 22	20.02 20.03 20.03	22.0	10.0	10.5	17.5	0 00	400	11.9	23.0	27.4	+ L	000
Olivia / Linner / Press	Pensile	11.7° F	8.3 1.0	0.00 6.70 to	3.3	1.0	10.0	7 9		30.1	0.4	13.0	0.70	1.6
	1 1	37° F 115°	ڙ وه	 	0.3		00	. :-		8.000		8.0	***********	-
	Durtility at	770 1	0.8 8.6	-1.	6.001	580 ++4	0 20		3	6.7	000	0.3	ŧ. 9	eo
	1:	115° F	<b>£</b> ≁88	204	4	45	255		- Page	-25-	· #	25.05	3 %	- -#
1	3	lactor		82.82 8.4.	28 3		28.6	51.4+	25.1	36.9	24.8	8:1:8	33.5	29.7
	at		### 288 288	76.6 87.9	80.4	103+	103+ 63.0	103+	448	57.5 03+	103+	*++	8.08	0.1.0
	Consistency at		103.5 +80.5 +80.5 +80.5	84.7 40.1	41.2	72.1	103+ 25.5 25.6	8.09	26.3		22.3 22.9		818	2
	Come		27.3 20.3 11.7	27.72	2 : :	12.4	54.2 9.4 10.4	13.2	14.8		13.0		8.0	
1	300 %		000		25 to 31 30		13	-	866		60 0		#1	
1	Penetration at F 77º F		ಇ೦ಐಂ	aj∞ o	100 to 120 110 to 120	35 to 45	2019 100	5 5	2,23	0 34	52 Kg 75 Kg	0 1	27	-
	Per 115° F		2301	SE 2	Too soft 10	24 250	58 46	61	34 20 34	103	14 46 58	201	33	
	Material	4 T. D. T.	BPL cannel Barrett pitch IBM pipe coating	IBM No. 2.			JM pitch No. 2. JM asphalt. Barber plastic pipe coating	Robertson 998	National saturant Robertson 230 Floating (Pobes)	Resistor enamel	Lyne-Kote hot. Texas asphalt No. 30 St. N. J. brick filer. Bitnmostio VXT	Bitumastic	Floatine (Milwaukee). East Ohio asphalt.	a Characteristics of original aembolt of contract

a Characteristics of original asphalt as given by the manufacturer.

#### 3. SOILS AT TEST SITES

For a complete understanding of the results of the National Bureau of Standards tests, detailed information as to the conditions to which the materials were exposed is essential. Descriptions of the profiles of the test sites are therefore presented in Appendix 2.

These descriptions were prepared in most instances by soil scientists of the U. S. Department of Agriculture. In a few cases the description of the site was prepared by the man who buried the specimens. The reader will see that in many cases the test trench was not uniform in all respects throughout its length. This contributed to the dispersion of the data. Under the conditions of the test, it was impracticable to secure completely uniform sites. However, the uniformity is probably greater than that encountered by pipe lines, and the lack of uniformity calls attention to the variability in corrosion which must be expected on pipe lines. From a practical viewpoint it may be fortunate that completely uniform sites were not available, as such sites might have obscured an important characteristic of underground corrosion.

The National Bureau of Standards has conducted or supervised corrosion tests at 128 locations scattered throughout the United States. In some instances, however, two or three tests of different materials have been conducted in the same type of soil. For this reason the Bureau's tests furnish information directly on only about 95 types of soils. Table 19 shows the numbers, types, locations, and other data relating to the soils in the National Bureau of Standards tests.

Table 19.—Test sites, soil numbers, types, locations, and other information relating to soils at NBS, AGA, and API test sites.

[G, good; F, fair; P, poor; VP, very poor]

Soila	Soil type	Location	Mean temper- ture ^b	Annual precipi- tation ^b	Interna drain- age of test site
	NATIONAL BUF	REAU OF STANDARDS SIT	res		
1 2 3 4 5	Allis silt loam. Bell clay Cecil clay loam. Chester loam Dublin clay adobe.	Cleveland, Ohio. Dallas, Tex. Atlanta, Ga. Jenkintown, Pa. Oakland, Calif	°F 49.2 65.5 61.2 54 56.4	Inches 33.8 36.2 48.3 40 23.4	PPOFP
6 7 8 9	Everett gravelly sandy loam. Maddox sit loam. Fargo clay loam. Genesee sitt loam. Gloucester sandy loam.	Seattle, Wash. Cincinnati, Ohio Fargo, N. Dak. Sidney, Ohio. Middleboro, Mass.	51.0 53.2 39 51.2 50	34.0 38.6 21 39.0 41	G F P P F
11 12 13 14 15	Hagerstown loam. Hanford fine sandy loam. Hanford very fine sandy loam. Hempsted silt loam. Houston black clay.	Baltimore, Md. Los Angoles, Calif. Baltershold, Calif Paul, Mire San Agrotao, Tex.	55.4 62.4 64.6 44.2 68.9	42.6 15.2 5.6 27.2 27.2	G F F 1
16 17 18 19 20	Kalmia fine sandy loam Keyport loam. Knox sitt loam. Lindley silt loam. Mahoning silt loam.	Mobile, Ala. Alexandria, Va. Omaha, Nebr. Des Moines, Iowa Cleveland, Ohio.	67.3 55 50.6 49.5 49.2	61.6 42 27.8 32.0 33.8	F G G
21 22 23 24 25	Marshall silt loam Memphis silt loam Merced silt loam Merrimac gravelly sandy loam Miami clay loam	Kansas City, Mo. Memphis, Tenn. But opwiew Cart. Notatos Miss. Mily mae, Wi	54.4 61.6 65 50 46.1	37.1 47.7 6 41 30.1	F G F G F
26 27 28 29	Miami silt loam Miller clay Montezuma clay adobe Muck Muscatine silt loam	Springfield, Ohio Bunkie, La. San Diego, Calif. New Orleans, La. Davenport, Iowa	53 67 61.0 69.3 49.9	37 56 10.3 57.4 32.1	(† P VP P
31 32 33 34 35	Norfolk fine sand Ontario loam Peat Penn silt loam Ramona loam	Jacksonville, Fla. Rochester, N. Y. Milwaukee, Wis. Norristown, Pa. Los Angeles, Calif.	69.3 47.6 46.1 54 62.4	47.4 32.8 30.1 40 15.2	G VP F G
36 37 38 39 40	Ruston sandy loam St. Johns fine sand Sassafras gravelly sandy loam. Sassafras silt loam. Sharkey clay.	Meridian, Miss. Jacksonville, Fla. Camden, N. J. Wilmington, Del. New Orlcans, La.	64.0 69.3 54 51 69.3	53.0 47.4 40 40 57.4	G P G F P
41 42 43 44 45	Summit silt loam Susquehanna clay Tidal marsh Wabash silt loam Unidentified alkali soil	Kansas City, Mo. Meridian, Miss. Elizabeth, N. J. Omaha, Nebr. Casper, Wyo.	54.4 64.0 52 50.6 47.2	37.1 53.0 43 27.8 15.3	F P VP G P
46 47 51 52 53	Unidentified sandy loam Unidentified silt loam Acadia clay Lake Charles clay loam Cecil clay loam	Denver, Colo Salt Lake City, Utah Spindletop, Tex League City, Tex Atlanta, Ga.	50.0 51.6 69 69 61.2	14.1 16.1 49 47 48.3	G P P G
54 55 56 57 58	Fairmount silt loam. Hagerstown loam. Lake Charles clay. Merced clay adobe. Muck.	Cincinnati, Ohio Baltimore, Md El Vista, Tex. Tranquillity, Calif. New Orleans, La.	53.2 55.4 69 63 69.3	38.6 42.6 49 8 57.4	P G P P VP

(See footnote at end of table.)

Table 19.—Test sites, soil numbers, types, locations, and other information relating to soils at NBS, AGA, and API test sites—Continued.
[G, good; F, fair; P, poor; VP, very poor]

		, , , , , , , , , , , , , , , , , , , ,			
Soila	Soil type	Location	Mean temper- ture ^b	Annual precipi- tation ^b	Internal drain- age of test site
	NATIONAL BUI	REAU OF STANDARDS SIT	res		
59 60 61 62	Carlisle muck. Rifle peat. Sharkey clay Susquehanna clay Tidal marsh	Kalamazoo, Mich. Plymouth, Ohio. New Orleans, La. Meridian, Miss. Charleston, S. C.	°F' 49 49 69.3 64.0 66.0	Inches 31 37 57.4 53.0 45.2	VP VP P P VP
64 65 66 67 68	Docas clay. Chino silt loam. Mohave fine gravelly loam Cinders. Gila clay	Cholame Flats, Calif. Wilmington, Calif Phoenix, Ariz. Milwaukee, Wis. Phoenix, Ariz.	58 62.4 69.7 46.1 69.7	16 15.2 7.8 30.1 7.8	VP F G VP F
69 70 71 72 76	Houghton muck. Me real V - open Mahorung sir orm Trumbu - oliy toam Cinno sit loam	Kalamazoo, Mich. Button willow, Calif. Austintown Junction, Ohio Yale, Ohio Los Angeles, Calif.	49 65 62.4	31 6  15.2	P P VP VP F
77 78 101 102	Susquehanna clay . Caddo fine sandy loam . Billings sift loam (low alkali) . Billings sift loam (modernis alant) . Billings sift loam . (modernis alant) .	Meridian, Miss Latex, La Grand Junction, Colo do	64.0 64.0 52.0 52.0	53.0 53.0 8.8 8.8	P VP F F
103	(Light Stall)	qo	52.0	8.8	F
104 105 106 107	Cecil clay	Charlotte, N. C	60.2 64.2 60 60.1 61.2	46 1 44.5 46 46.3 48.3	00000
109	Fresno fine sandy loam	Fresno, Calif	63	9	P
110	(low alkali) Fresno fine sandy loam	do	63	9	P
111	(moderate alkali) Fresno fine sandy loam	Kernell, Calif	63	9	P
112 113	(high alkali) Imperial clay (moderate alkali). Imperial clay (high alkali)	Niland, Califdo	71 71	2 2	F F
114 115 116 117 118	Lake Charles clay Memphis silt loam Merced clay Merced clay loam adobe. Niland gravelly sand (low alkali)	El Vista, Tex. Vicksburg, Miss. Los Banos, Calif. Tranquillity, Calif. Niland, Calif.	69 65.6 63.4 63 71	49 51.9 8.1 8	PGFPF
119 120 121 122 123	Norfolk sandy loam Norfolk sand do Panoche clay loam Susquehanna clay	Macon, Ga. Pensacola, Fla. Tampa, Fla. Mendota, Calif. Shreveport, La.	64.2 67.7 71.8 63 65.8	44.5 57.9 49.4 6.4 43.4	<del>ዕ</del> ዕሪፍ ዮ
124 125	Susquehanna silt loam Susquehanna fine sandy loam	Troup, TexShreveport, La	66.0 65.8	42.7 43.4	P
	AMERICAN (	GAS ASSOCIATION SITES	,		
1 2 3 4 5	Cinders	Pittsburgh, Pa. Milwaukee, Wis. Brockton, Mass. Atlantic City, N. J. West Palm Beach, Fla.	46.1	30.1	VP P P P
6 7 8 9 10	. do Cecil clay loam do. Susquehanna clay Miller clay do	Miami, Fla. Atlanta, Ga. Raleigh, N. C. Shreveport, La. do. Bryan, Tex.	61.2 60.1 65.8 65.8	48.3 46.3 43.4 43.4	PGGPPP

(See footnote at end of table.)

Table 19 .- Test sites, soil numbers, types, locations, and other information relating to soils at NBS, AGA, and API test sites -Continued.

[G. good: F, fair: P, poor; VP, very poor]

Soila	Soil type	Location	Mean temper- ture ^b	Annual precipi- tation ^b	Internal drain- age of test site					
	AMERICAN	GAS ASSOCIATION SITES								
12. 13. 14.	White alkali soil Bluck i lkall soil. Marsher shiere i	Los Angeles, Califdo Kansas City, Mo	°F 62.4 62.4 54.4	Inches 15.2 15.2 37.1	G G					
AMERICAN PETROLEUM INSTITUTE SITES										
II. III. IV. V.	Bell clay	Temple, Tex. Arkansas City, Kans. Beaumont, Tex League City, Tex Preble, Ind	69	49	P VP					
VII. VIII. VIII. IX. X.	Not determined. Oswego silf-loam Andic chi Harror tha soudy loam Musecono si then	Council Hill, Okla	69	49						
XI. XII. XIII. XIV.	Not determined Merced clay loam Miller clay Hagerstown silt loam	Skiatook, Okla Mendota, Calif Bunkie, La Chambersburg, Pa	63 67	6.4 56	P VP					
άvi.	Docas clay	Cholame, Calif.	58	16	VP					

a Different numbers for soils that appear to be identical indicate either different sites in the same locality or that different classes of materials with the internal office of the classes of materials with the classes of the decimal of the fundation of the decimal point are for some nearby city.

The soil types of soils 71 to 78 are subject to change upon better identification.

The selection of the test sites was governed to a large extent by the distribution of pipe lines and pipe networks, that is by the importance of the soil with respect to underground construction. Each test site represents an important soil condition, but as within an area of a few square miles several quite different soils usually exist, they are not necessarily representative of the soil conditions prevailing in the region of the site. This is illustrated by figure 7, which shows the soil series in a 6- by 7½-mile area just east of Baltimore, Md. [46].

It should be obvious that a test of materials in any one locality may not indicate the behavior of the materials in a different soil which may lie within a short distance from the test site. Moreover, as the soil horizons of a single soil may differ widely in physical and chemical properties, degree of aeration, and moisture content, a test of pipe materials in one soil horizon may yield results different from those of a test of the same materials exposed to another horizon of the same soil type. For example, the averages of the two deepest pits on five ferrous materials buried for 5 years in Rifle peat near Plymouth, Ohio, were 38, 37, 21, 24, and 67 mils, respectively. The corresponding values for the pit depths on the same materials exposed for 7 years at the same site were 30, 34, 16, 17, and 62 mils, respectively. An examination of the site showed that the two sets of specimens were placed in two paralleled trenches only a few feet apart. The depths of the trenches were approximately the same, but the 5-year-old specimens were laid in the peat horizon, whereas the 7-year-old specimens were placed in the clay subsoil just below the peat. The surface of the ground was level, but the thickness of the peat layer varied.

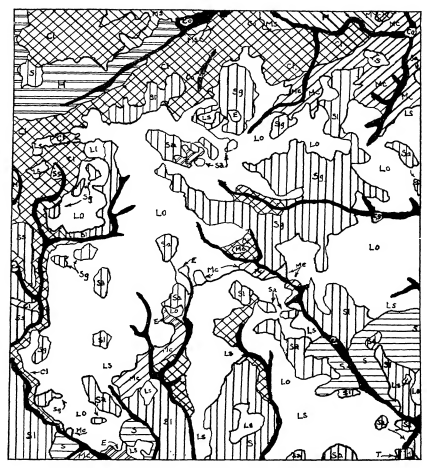


FIGURE 7.—Soil series in a 6- by 71/2-mile area near Baltimore, Md.

	LLD 1. DOUB GOT FOR THE CO	og 1/2 mile area mear Dairemer, and
C1	= Chester loam.	Mc = Montalto clay loam.
	= Congaree silt loam.	S = Susquehanna silt loam.
	= Elkton silt loam.	Sa = Sassafras sandy loam.
	= Hagerstown loam.	Sg = Sassafras gravelly loam.
	= Iredell silt loam.	SI = Sassafras loam.
	= Leonardstown loam.	Ss = Sassafras silt loam.
	= Leonardstown silt loam.	T = Tidal Marsh.
M	= Manor loam.	

For convenience, the descriptions of the soils at the Bureau sites have been arranged according to the arbitrarily assigned numbers of the test sites. Those who are interested in the classifying of the soils as to the great soil groups to which they belong should refer to table 1.

Tables 20 to 23 show the more important properties of the soils to which specimens were exposed. Since these data were obtained, it has been discovered that the pH of some poorly drained and poorly aerated soils changed after they were exposed to air [47]. As the soils were

air-dried and pulverized before the analyses given in table 20 were made, it is probable that the pH values given for some of the soils are incorrect. New values will be obtained as soon as conditions permit.

Table 20.—Chemical properties of soils at National Bureau of Standards test sites.
[A, Alkaline; C, not determined.]

***************************************			[A, Alka	line; C, r	not deter	minea.				
Soilb		Total acidity,	Resis-	Compo	sition of	water ex	tract, m	g-eq per	100 g of a	soil
No.	pH	mg-eq per 100 g of soil	tivity at 60° F	Na+K as Na	Ca	Mg	CO ₃	HCO3	Cl	SO ₄
1 2 3	7.0 7.3 5.2	$11.4 \\ 3.5 \\ 11.5$	Ohm-cm 1,215 684 30,000	0.72	0.25 1.09	0.43	0.00	0.09 1.18	0.09	0.83
4 5	5.6	7.6 6.5	6,670 $1,346$	93	0.48	.10		0.69	.03	
6 7 8 9	5.9 4.4 7.6 6.8 6.6	12.8 29.8 A 7.2 3.6	45,100 2,120 350 2,820 7,460	1.42	1.72	2.55	.00	.71		4.43
11 12 13 14	$\begin{array}{c c} 7.1 \\ 9.5 \\ 6.2 \end{array}$	10.8 2.5 A 5.6 5.0	11,000 3,190 290 3,520 489	0.39 6.23 2.18	0.50 .09	0.16 .13	.00		.00 1.64 0.13	0.14 3.76
16 17 18 19 20	4.5 7.3 4.6	11.8 19.1 1.4 10.9 1.5	8,290 5,980 1,410 1,970 2,870	0.27 .38 .25			.00			 .25 .46 .15
21 22 23 24 25	9.4	9.5 9.7 A 12.6 4.7	2,370 5,150 278 11,400 1,780	8.38	.38	.41	.02	1.87	i.i2 0.03	5.57 0.10
26	$\begin{array}{c c} 6.8 \\ 4.2 \\ 7.0 \end{array}$	2.6 3.7 C 28.1 2.6	2,980 570 408 1,270 1,300	.27 .53 1.50 2.15 0.32	.50 1.86 0.06 1.92 0.65	.31 1.12 0.18 1.55 0.40	.00 .00 .00 .00	$0.70 \\ 2.00 \\ 0.12 \\ .00 \\ .71$	.03 .08 .99 1.69 0.09	1.54 0.89 2.30 0.24
31 32 33 34	6.8	1.8 0.5 36.0 7.0 5.7	20,500 5,700 800 4,900 2,060		7.30 0.68	12 4.06 0.49	.00		0.00 0.00	42 2.13 0.35
36 37 38 39 40	4.5 5.6 6.0	4.6 15.3 1.7 6.6 9.4	11,200 11,200 38,600 7,440 970	0.56	0.58	0.44	.00	0.93	.07	.28
41 42 43 44 45	5.5 4.7 3.1 5.8 7.4	11.0 28.2 36.8 8.8 A	1,320 13,700 60 1,000 263	45.10 1.05 8.15	5.17 1.08 3.70	9.45 0.66 .70	.00	.78 .00 1.97 0.24	.04 43.30 0.82 .18	.46 37.00 0.41 11.98
46	. 7.6	C 3.0 13.2 A 9.6	1,500 1,770 190 234 17,794	0.67 10.27 4.20	0.72 15.55 .33		.00	.88 .56 1.36	.06 5.75 1.33	0.48 22.00 1.26
54 55 56 57 58	7.1 7.7	3.5 10.9 4.5 A 79.3	886 5,213 406 128 712	0.59 3.12 23.40 2.03	10.10 0.69 13.50 2.23	0.59 	.00	.70 .80 .34 .00	0.08 1.59 1.15 0.47	9.05 3.04 37.50 2.54
59 60 61 63	2.6 5.9 4.1	33.3 297.4 8.6 24.2 100.2	1,659 218 943 6,922 84	1.03 2.91 0.73	3.08 10.95 0.68	2.70 2.86 0.33	.00	.00 .00 .71	3.47 0.00 .10	1.04 56.70 0.91
64 65 66 67 68	8.3	A A A A	62 148 232 455	28.10 7.65 6.55 0.77	2.29 12.40 0.51 3.03	0.76 2.20 0.18 53	.00	.89 1.30 0.73 .55	28.80 6.05 2.77 0.08	0.26 16.90 2.97 2.89
		1			1		All I			

(See footnote at end of table.)

Table 20.—Chemical properties of soils at National Bureau of Standards test sites—Continued.

[A, Alkaline; C, not determined.]

[-1, 120 document]											
Soilb		Total acidity,	Resis-	Compo	sition of	water ex	tract, m	g-eq per	100 g of	soil	
No.	pН	mg-eq per 100 g of soil	at 60° F	Na+K as Na	Ca	Mg	CO ₃	HCO3	Cl	SO ₄	
69		33.3 A	Ohm-cm 1,659 278	1.03 8.38	3.08 0.38	2.70 0.22	.00	.00 1.87	3.47 1.12	1.04 5.57	
71 $72$ $73$	7.2		762			1					
74 75 76 77										::::::	
101 102 103 104 105	7.3 7.3 7.3 4.6 4.8	A A 11.0 12.9	261 103 81 8,500 28,000					0.66 .56 .18	1.56 4.67 11.09	22.48 36.82 25.70	
106 107 108 109	4.8 4.8 4.9 8.4 10.2	12.8 11.8 11.2 A A	25,000 54,400 44,400 497 531	2.62 3.53					1.56 0 79		
111 112 113 114	7.3 7.4 7.4 7.1 6.9	A A A C 4.7	51 149 102 320 3,450	41.55 22.18 9.56 2.65	16.21 14.09 0.84 0.68	.44 1.29 .51 0.26	0.00 .00 .00 .00	0.58 .36 .63 .77	34.58 10.94 6.26 1.84	23 41 25.98 4.06 0.93	
116 117 118 119	9.2 8.5 7.3 4.7 5.7	A A 9.7 1.3	320 106 273 10,800 34,400			.18 .83 .64		2.10 1.55 0.20	1.17 2.89 5.01	1.57 46.53 0.86	
121	4.8 7.4 4.1 4.4 3.9	1.2 A 21.9 28.1 28.3	1,160	2.32	0.05	.26	.00		0.07		

 $^{^{\}rm a}$  Measurements and determinations by I. A. Dension and R. B. Hobbs.  $^{\rm b}$  See table 19 for names and locations of soils.

	o de N
ない。 では、これでは、これでは、これでは、これでは、これでは、これでは、これでは、これ	Moisture equiva- lent
	Airpore
REVERNATAR REPRESE SAR SAR SAR REPER	Apparent specific gravity
	1.45
UK 대단도로 등록 등등로 등등로 등등로 갖지않는 그 건강 그 건강 등록 작용하고 있다고 등 본 옷 작용한	No.
対象対象	Moistare equiva- lent
a Baratus inang	Airporr
5 32-23-34 desc	Apparent specific gravity
10   10   10   10   10   10   10   10	Volume shrink- age

47	58388	82828	38222	57552	E80070	C1 42 C0 1C +	out. NO.	S. Hirs
8883	88888	88843	88888	±8386	88888	Inches 90 40 82 86 30	Беріл	7
9.2.30 2.4.1.0	25.11 6 6 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5	21.0 1.4 2.1 97.3 42.1	15.7 19.6 3.2 1.2 72.0	50.4 50.4 1.3	25.8 25.8	25 25 25 25 25 25 25 25 25 25 25 25 25 2	Sand	
56.7 24.1 44.9	55 ± 52 53 ± 55 55 55	#2:058 11:158	22.4 25.9 26.9	78.5 28.5 4.6 1.2 28.5 28.5 28.5	23.8 26.1 27.7 29.4 21.1	87.7 24.9 53.0	SII	Mechar
\$6.1.28 6.1.28	26.0 17.6 47.1	15 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8	52883 5898 6898	26 5 20 3 20 3	7 2 68 6 6 6 1 6	52.5 44 9 117 7 1	Clay	Mechanical analysis
35.0 40.9 25.8	82.54.119 82.78.88	21 s 71 s 26.1	223.7 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1 277.1	9.5 62.0 21.8 15.0	50.55 50.55 98.77 98.87	31:1 40 ± 1 9:9 30:2	Colloid	
24.7 45.1 5.5	22.2 17.0 24.9	24.8 25.7 15.9	15.9 87.7 26.0 38.7	25128 25128	12.2 41.2 20.2 16.4	38.0 19.1 4.4 40.8	percentage	G

There 99.— Understand mediges of soils of Indiana Berene of Standards test sites. Pages 21 - Physical Properties of soils at Arthough Darrow of Mondards and sole

Table 23.—Properties of the soils in the American Gas Association and American Petroleum Institute coating tests.

Soil No.	Soil type	Mois- ture equiva- lent	Volume shrink- age	Air- pore space	Ap- purent specific gravity	pH (aver- age)	Total acidity (mg-eq per 100 g of soil)	Resistivity, at 60° F
	AMERIC	AN GAS	ASSOCI.	ATION I	TEST SIT	168	,,	
1 2 3 4 5	Cindersdo	% 18.2 9.5 50.3 93.7 74.2	%  15.0	%		5.5 7.2 3.6 3.0 4.3	42.5 38.0 75.8	Ohm-cm 730 380 44 32 1,180
6 7 8 9	do. Cecil clay loamdo. Susquehanna clay Miller clay	29.1 35.0 37.6 35.8 32.0	40.4 38.8 35.5 37.2			5.7 5.8 6.9 4.6 7.4	$\begin{array}{c} 22.2 \\ 13.3 \\ 6.3 \\ 15.7 \\ 1.1 \end{array}$	1,650 43,800 16,000 6,840 870
11 12 13 14	do	47.8 35.6 16.6 31.0	49.9 45.0 14.0 41.0			7.2 7.3 9.2 6.5	0  12.8	1,000 93 1,700 3,150
	AMERICAN	PETRO	LEUM IN	STITUT	E TEST	SITES		
III. III. IV. V.	Bell clay Oswego silt loam Lake Charlos clay I nke Charlos clay loam Macri sil: I m:	$31.6 \\ 20.7 \\ 40.7 \\ 21.1 \\ 21.5$	28.0 $5.1$ $35.8$ $6.7$ $6.4$	5.1 $19.2$ $2.6$ $5.1$ $6.9$	2.00 $1.76$ $2.00$ $1.93$ $1.87$	8.4 $7.0$ $6.8$ $7.2$ $6.3$		947 1,295 495 1,485 2,201
VII. VIII. VIII. IX.	Unidentified Oswego silt loam Acadht clay Hanford fine sandy loam Muscatine silt loam	$25.4 \\ 20.6 \\ 47.6 \\ 9.8$	$\begin{array}{c} 11.6 \\ 7.0 \\ 42.7 \\ 0 \end{array}$	5,5 13,8 2,9 20,9	1.78 2.04 2.04 1.47	5.4 5.4 8.9 6. +		5,180 3,510 259 353
XI. XII. XIII. XIV. XVI.	Unidentified Merced clay loam Miller clay Hagerstown silt loam Docas clay	15.7 $36.8$ $38.0$ $21.3$ $36.8$	$\begin{array}{c} 0 \\ 31.2 \\ 30.2 \\ 1.4 \\ 29.1 \end{array}$	16.6 7.2 2.5 8.3 3.8	1.65 1.84 2.01 1.46 1.88	$\frac{5.2}{8.0}$ $\frac{7.0}{6.4}$ $\frac{6.4}{8.4}$		440 61 674 5,088 155

### 4. FACTORS AFFECTING RESULTS OF CORROSION TESTS

Among the factors that affect the results of corrosion tests are depth of burial, mill and foundry scale, diameter of the pipe, area of pipe surface, and duration of exposure. In order to gain an idea of the nature and relative magnitudes of the influence of these factors, several supplementary tests were carried out.

#### (a) DEPTH OF BURIAL

In general, aeration in soils will decrease as the distance below the surface increases and, therefore, the corrosivity of the soil should be expected to increase with depth. In fact, pipe-line operators have often reported more severe corrosion where their lines are deeper than usual. In order to investigate the effect of depth of burial, samples of 3-inch steel pipe were buried at different depths in seven soils having different general characteristics. The results of this series of tests are given in table 24. In five of the seven soils, the amount of corrosion was greater at the greater depths. In two of the soils, however, the reverse was true-This is probably due to the counteracting effect of the subsoils. For instance, in the case of St. John's sand (soil 37), in which corrosion was greater near the surface there is an impervious hardpan layer about 6

inches thick at about 18 inches below the surface of the ground. If pipes are laid in or above this layer, they may be surrounded by water much of the time because the water can not get through the layer. If pipes are laid deeper, they will be placed in sand, which will let the water pass more readily. In the arid regions of the West, the concentration of the alkali may be either near the surface or at a considerable depth, depending on the prevailing direction of the movement of moisture in the soil. Thus it can be seen that, although corrosion in general increases with depth of burial, other influences, such as drainage and the characteristics of soil horizons, may be sufficient to mask completely the effect of depth.

Table 24.—Effect of depth of burial on corrosion of 3-inch steel pipe.

[Average of four specimens]

Soil number	Type of soil	Depth of burial	Loss of weight	Maximum penetration
	EXPOSURE OF 11.70	YEARS		
37 37 37	St. John's fine sanddodododododo	in. 12 24 36 48	$0z/ft^2$ $10.13$ $9.05$ $5.27$ $4.71$	Mils 67 64 50 44
	EXPOSURE OF 9 27	YEARS		
103 103 103	Billings silt loam (high alkali) do do do	14 22 28 48	16.33 17.99 17.66 20.76	118 136 148 172
	EXPOSURE 11.71	YEARS		
104 104 104 104	Cecil clay do do do do do do do do do	18 30 36 52	5.61 8.18 9.07 7.92	80 102 110 166
	EXPOSURE 11.70	YEARS		
107 107 107	Cecil fine sandy loamdododo	12 24 36	3.89 4.40 4.85	53 64 89
	EXPOSURE 9.24 Y	EARS		
110 110 110	Fresno fine sandy loamdodo	19 26 42	13.96 18.18 18.63	113 124 159
	EXPOSURE 5.93 Y	EARS		
113 113 113 113	Imperial clay (high alkali)	12 22 28 40	17.90 20.05 21.66 21.69	166 211 212 198
	EXPOSURE 9.27 Y	EARS		
117 117 117	Merced clay adobedododododododo	12 24 36 48	20.28 19.09 19.70 12.03	127 122 87 83

#### (b) MILL AND FOUNDRY SCALE

The oxide which forms on the surface of a pipe as the result of rolling or casting is at a more noble potential than that of the unoxidized metal. and the oxide probably corrodes more slowly. Indeed, the corrosionresisting properties of stainless steel are attributed to a very thin oxide film which reforms whenever it is removed. A continuous oxide film should retard or prevent corrosion. However, if unoxidized metal is exposed adjacent to the oxide, a difference of potential is created which tends to accelerate the corrosion of the unoxidized metal. Unfortunately, the oxide scale on a pipe is seldom continuous. In 1926, specimens of several materials in their original condition and similar materials with the mill scale removed were buried in seven soils. Table 25 shows the rates of corrosion of these specimens after approximately 8 years of exposure [45]. To facilitate the comparison of the materials with and without mill scale, the ratio of the number of cases in which the scalefree material was superior to the scale-coated specimens is given at the bottom of each part of the table. It is evident from the table that the majority of the specimens from which the scale had been removed corroded somewhat less than those having a coating of mill scale, that is, mill and foundry scale did not afford any protection to the specimens. However, the improvement is too slight to warrant the removal of mill scale.

Table 25.—Effect of mill and foundry scale on corrosion.

[Period of exposure was approximately 8 years]

		Pit ca	Pit cast iron		delayand east iron		eel	Wrou	tht iron
Soil	Soil type	Nor- mal, A	Ma- chined MC	Nor- mal,	Ma- chined, MD	Nor- mal, P	Ground and pol- ished, M	Nor- mal, K	Rough ground, D
	RATI	S OF LO	OSS OF	WEIGH	T (oz/f	l ³ )/yr	·		
13 24 28 29 42 43 45	Hanford very fine sandy loam. Merrimae gravelly sandy loam. Montezuma elay adobe. Muck. Susquehanna elay. Tidal marsh. Unidentified alkali. Ratioa.	1.12 0.11 3.94 2.31 0.70 1.08 3.50	1.29 0.13 0.09 0.43 1.00	0.05 .17 1.14 5.28 0.59 3.08 3.55	1.10 0.16 1.27 0.47 1.20 2.59 5:6	1.70 	1.51 	1.12 0.22 1.18 0.88 0.88 .67 2.51	1.01 0.18 22 0.80 .67 2.17 4:6
	RA	TE OF	PENET	RATION	V (mils/	yr)			
13 24 28 29 42 43 45	Ratio ^a	17.8 3.6 20.4 11.4 17.9 14.1 22.3	12.8 3.2 5.0 15.9 5.2 5:5	7.8 3.8 6.8 10.7 7.0 27.5 14.3	9.1 2.4 8.9 8.9 11.5 15.7	19.3  10.5 8.8 10.8 16.6	15.3  9.8 10.0 8.1 13.0 4:5	11.9 4.7 	11.1 3.8 

Ratio of cases of superiority to total number of cases of comparison.

## (c) DIAMETER OF PIPE

When tests of small sections of pipes are planned, the question of the effect of the diameter of the pipe may arise. If the diameter of the pipe

is so large that the pipe lies in two soil horizons, or if only part of it is under water, or if there is a considerable difference in the distances from the surface of the ground to the top and bottom of the pipe, differential aeration circuits will be set up which will affect the distribution of the corrosion on the surfaces of the pipe and possibly the amount of corrosion also. It is not often, however, that such large diameter pipes are used in corrosion tests.

Table 26 shows the effect of diameter on the depths of the maximum pits on two different materials as calculated in three different ways. From each soil there were removed one specimen of 6-inch cast-iron pipe, two specimens of 3-inch cast iron, and two specimens of 3-inch steel pipe, all 6 inches long. There were also removed from each soil two specimens of cast iron 12 inches long with external diameters of approximately 1.9 inches, and two 10-inch lengths of 2-inch open-hearth steel. By treating the two specimens of each material, except the 6-inch cast iron, as a single specimen having twice the area of one, it is possible to compare the pit depths on specimens of different materials and diameters having approximately the same areas. Columns 1 to 5 of table 26 permit such a comparison.

Table 26.—Average of maximum pit depth on pipes of different diameters for four periods of exposure.

		Single	maxim	um pit		A	verage o	of two d	eepest p	oits	Weig	ghtedb n pit d	maxi-
Soil num- ber		Cast iro	on	Ste	eel		Cast ire	on	Ste	eel	of	cast in	on
	6 in.	3 in.	1¼ in.	3 in.	2 in.	6 in.	3 in.	1¼ in.	3 in.	2 in.	6 in.	3 in.	11/4 in.
	1	2	3	4	5	6	7	8	9	10	11	12	13
*51 53 55 56 58	Mils 169 68 37 139 129	Mils 151 72 48 147 125	Mils b212+ 61 93 166+ 195+	Mils 60 58 69 156+ 70	Mils 136+ 53 60 106+ 97	Mils 155 63 33 125 124	Mils 128 58 42 133 113	Mils 206 + 55 87 166 + 194 +	Mils 52 55 64 150+	Mils 124 + 50 55 95 + 75	Mils 145 54 31 111 114	Mils 101 49 38 123 105	Mils 201+ 52 82 160+ 134+
a59 60 61 62 63	50 °105s 60 82 43	43 111 67 79 42	58 102+ 74 103 81	39 114+ 56 70 134+	39 36 76 85 51	47 100s 54 77 38	37 100 58 71 38	35 81 + 68 92 69	35 88+ 50 58 98+	25 27 63 74 41	43 89s 49 75 32	36 96 54 65 34	91+ 63+ 58 75 66
64 65 66 67	225 140 168 141s	195 162 167 164	163+ 123 146+ 231+	178+ 120+ 171 139s	150 + 83 133 + 154 +	216 132 157 122s	187 143 164 142	155+ 101 119+ 217+	160 102+ 154 112	148+ 79 132+ 147+	197 121 143 110s	195 129 150 118	137+ 96 103 200+
Aver- age	111	112	129	102	90	103	101	118	89	81	94	92	108

Average for 3 periods of exposure only.
 A plus (+) sign indicates that 1 or both removals were punctured during 1 or more periods.
 s indicates that uniform corrosion occurred on 1 or more specimens.

In 7 of the 14 soils shown in table 26, the 6-inch cast-iron specimens developed shallower maximum pits than were observed on equal areas of 3-inch or 1½-inch cast specimens, and the average of the maximum pits on the 6-inch specimens is less than the averages for either of the other two sizes of cast-iron pipes. The 3-inch cast specimens developed shallower maximum pits than were found on the 1½-inch cast specimens in 6 of the 14 soils, and the average of the maximum pits in the 14 soils is slightly greater for the 1½-inch cast iron. The 3-inch steel specimens

developed shallower maximum pits than were observed on the 2-inch steel specimens in 5 of the 13 soils for which comparisons are possible. A comparison of specimens of different diameters on the basis of the averages of the two deepest pits yield similar results (columns 6 to 10).

In earlier reports, the pit depths were weighted to take account of the difference in the areas of the specimens. The depths of the four deepest pits were averaged to obtain the recorded maximum pit depths on the 6-inch cast-iron specimens, whereas the depths of the two deepest pits on each of two 3-inch specimens were averaged to obtain the recorded maximum pit depth on the 3-inch specimens. The 1937 data [48] were treated in this way to obtain the values in columns 11, 12, and 13 of table 26. In six soils the 6-inch specimens showed deeper weighted maximum pit depths than the 3-inch cast-iron specimens, whereas in six other soils the weighted maximum pit depths on the 6-inch specimens were shallower. The averages of the weighted maximum pit depths for the 3 sizes of cast pipe in 13 soils differ by only 5 mils. It appears, therefore, that for small-diameter pipe, when the areas of the specimens are the same, the diameter has practically no effect on pit depth.

### (d) AREA OF PIPE SURFACE

Early in the course of the soil-corrosion investigation it was observed that there was a tendency for the specimens 3 inches in diameter to contain deeper pits than specimens of similar materials 1½ inches in diameter. This is illustrated in table 27, which permits a comparison of the maximum pit depths on areas of 66 and 126 square inches of pipe surfaces for two metals exposed for approximately 12 years in 38 soils. As the specimens differed in diameter, as well as in area, it is possible that the difference in pit depths is affected by differences in curvature of the specimens. However, according to Shepard [49] the greater curvature of the smaller specimens should tend to concentrate the corrosion and consequently intensify the pitting on the smaller area.

The apparent effect of area seems to be different for different soils and to be slightly different for the two materials. Data, as well as theoretical considerations, indicate that the pit-depth-area relation is influenced by soil characteristics, although just what the characteristics are has not been determined. Sufficient data are not available to determine whether or not the relation is the same for all ferrous materials, but data on pitting factors suggest that the corrosion of wrought iron is somewhat more uniform and that of pit-cast iron somewhat less uniform than that of steel. The difference, however, if any exists, is not great.

The observed values of maximum pit depths obtained from measurements made on a 12-inch oil line in Miller clay [35] are shown in table 28 as a function of the area inspected. In this table are tabulated the depths of the deepest pits on each of a number of 20-foot lengths of pipe and also the averages for the depths of the deepest pits on each foot of the corresponding lengths. The data for individual lengths are given in order to show how much pit depths may vary in a single type of soil. It will be noted that the deepest pit on 48 lengths of pipe, 950 feet, was 306 mils; the average for depths of the deepest pit on each length of pipe, 20 feet, 208 mils, and the average of the deepest pits on each foot of pipe, 126 mils,

Table 27.—Effect of area inspected on the observed maximum pit depths on National Bureau of Standards specimens.

[Pit depths in mils.]

Soil No.	7	epth of m			0.337	-			<del></del>
Y .		epth of m	aximum	pre	Soil No.		epth of m	aximum	pit
	Besserr	ner steel	Wroug	ht iron		Bessen	ner steel	Wrou	ght iron
Area→	a66 in.2	^b 126 in. ²	66 in.²	126 in.²	Area→	66 in.2	126 in. ²	66 in.2	126 in.º
1 3 4 5	119 71 85 49	101 72 84 55	81 90 88 52	83 96 93 46	27 30 31	73 61 36 48	83 69 45 70	52 52 31 47	59 56 45 67
6 7 8 9	19 46 77 36 43	40 65 119 50 60	22 43 103 46 54	38 48 84 47 52	33 34 35 33 37	92 42 28 54 79	103 60 90 47 99	108 42 42 49 73	111 66 59 50 82
12. 15. 16. 17.	57 58 61 42 44	72 85 69 47 50	58 61 67 40 48	91 83 60 42 43	38 39	37 53 73 87 131	52 113 83 80 97	44 58 71 127 95	29 74 70 89 98
19	78 78 71 25 48	61 56 71 31 39	71 52 66 25 47	79 51 60 29 45	43 44 45 46	108 63 113 75 30	90 99 137 136 29	80 58 106 72 59	173 73 92 80 38
					Average for 33 soils.	63	74	63	68

a Specimens 113 in. in diameter. b Specimens 3 in. in diameter.

TABLE 28.—Relation between area inspected and the observed depth of the deepest pit.

Pipe section number	Maximum pit depth on each lengtha	Average pit depth for 1-ft. lengths	Pipe section number	Maximum pit depth on each length	Average pit depth for 1-ft lengths
12334456	Mils 176 126 183 205 208 202	Mils 90 8 58.6 125.0 126.2 136.7 116.7	25. 26. 27. 28. 29.	Mils 287 185 122 290 258 160	Mils 180.0 101.2 65.7 154.7 126.3 94.5
7	227 174 207 217 137 257	122 0 138.3 144 3 105.3 79.6 159.3	31	125 258 233 190 188 207	67.9 144 4 138.5 115.9 115.8 127.4
3	^b 306 202 216 187 179 195	169.7 149.9 158.6 139.8 91.3 127.9	37. 38. 39. 40. 41. 42.	253 253 154 251 274 185	154.6 130.6 94.9 97.6 133.7 121.1
9	200 195 206 245 247 240	140.3 143.2 127.7 147.9 132.4 161.1	43. 14. 45. 46. 47. 48.	192 164 195 195 237 212	138 0 95.5 123.4 141.0 142.8 138.2
			Average	208	126

a Approximately 20 ft of 12-in. pipe, area 67 ft2.

The relation between the average of the depths of the deepest pits on unit areas and the size of that unit is illustrated in figure 8. The data

^b Deepest pit on 950 ft of pipe.

for this figure are measurements of the deepest pit on each 20-foot length of 15 miles of a 10-inch pipe line traversing a sandy-loam soil. From these data it was possible to determine the maximum pit depth on areas corresponding to any number of lengths of pipe up to the length of the line.

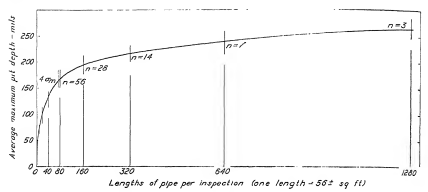


Figure 8.—Relation of maximum depth of pits to area inspected. n = number of inspections. The short vertical lines, 4  $\sigma_m$  indicates the probable range of variations in the data.

The lengths of the vertical lines crossing the curve represent four times the standard error of the average of the maximum pit depths. The numbers on the line show the number of observations upon which the average was based. If the pit-depth-area relation were determined for a line traversing more than one soil, the data would obviously be affected by the variety of soils traversed.

Scott [50], after an extensive examination of the Bureau's soil-corrosion data and of the measurements of pits on areas up to 10,000 square feet on oil and gas lines, found that the relation between the maximum pit depth and the area from which it was selected could be represented approximately by the equation  $P = bA^a$ , in which P is the maximum pit depth associated with an area, A, and a and b are constants which differ for different conditions.

From an unpublished study of data from eight National Bureau of Standards test sites and from seven sets of pipe-line data, Ewing found that on the average, the equation  $P = P_1$  ( $C \log A + 1$ ), in which P is the maximum pit depth associated with an area A,  $P_1$  the maximum pit depth on a unit area, and C is a constant varying with conditions, fitted

the data better than did Scott's equation.

The fact that buried metal does not corrode uniformly can be accounted for by assuming that conditions over the surface of the metal are not uniform. This condition may be the result of lack of uniformity in the metal, differences in the soil in contact with the metal at different points, or differences in aeration resulting from the way the soil was placed over the surface of the metal. An explanation of the pit-deptharea phenomenon applicable only to small isolated specimens differing in area, is as follows: If a difference of potential exists between two areas on the surface of a specimen, the current which flows between these areas when the circuit is completed through an electrolyte is influ-

enced by the area of the cathode because of its effect on the resistance of the circuit and because of its effect on polarization. The longer the specimen, the larger will be the possible area of the cathode. However, there are few data that show definitely what is the area of the cathode associated with a single pit.

If the explanations that have been given fully account for the pitdepth-area relation, it might be expected that when the maximum pit depths on a number of specimens having large areas are averaged, some area would be found such that larger areas would not contain deeper pits.

Figure 8 suggests that such may not be the case, as the maximum pit appears to increase in depth with increase in the area chosen for the unit of inspection up to 15 miles of pipe or up to an area of approximately 72,000 square feet.

It seems possible that the apparent relation between pit depth and area arises, in part at least, from the use of the maximum pit depth as the criterion for corrosivity. The probability that a deeper pit will develop on a large area than on a small area must be a function of the relative sizes of the areas and must always be greater than 50 percent. It seems likely, therefore, that the pit-depth-area relation may be, in part at least, a probability relation, but that it is modified to some extent by certain factors related to soil characteristics. However the relationship may be regarded, it must be taken into account if soil-corrosion data are to be correctly interpreted.

### (e) DURATION OF EXPOSURE

The depth of the deepest pit usually is not proportional to the period of exposure of the metal to the soil. This is illustrated by figure 9. A number of empirical equations have been offered to express the relation of pit depth to the duration of the exposure. Fetherstonhaugh [51] suggested the equation  $D = A^3 \sqrt{T}$ , in which D is the depth of the pit at any time, T, and A is the pit depth for unit time of exposure. Put-

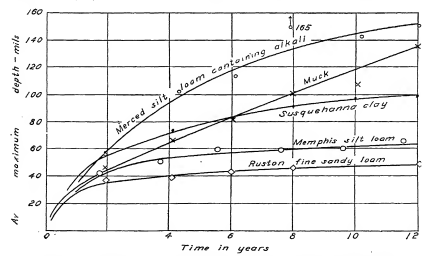


FIGURE 9.—Relation of maximum depth of pits to duration of exposure.

nam [52] proposed the equation  $P = KT^{0.61}$ , which differs from Fetherstonhaugh's equation only in the value of the exponent of T. These equations indicate that the change in pit depths with time follows a similar course in all soils.

Brennan [53], from a study of pipe-line data, decided that the relation of pit depth to time could be represented by the equation  $y = (A + BZ) \log (x/h + 1)$ , where y is the depth of the pit at a time x, Z is the Corfield soil-corrosivity index [54], and A, B, and h are constants to be derived from a study of test data.

Scott [55], after analyzing the National Bureau of Standards data, concluded that the relation of pit depth to time could be represented approximately by the equation P = UT/(B+T), in which P is the pit depth at time T, and U and B are constants which characterize the soil.

Ewing [35] found that the occurrence of leaks on pipe lines was consistent with the equation  $P = kT^n$ , in which P is the depth of the deepest pit (on equivalent areas) at time T, and k and n are constants depending on the characteristics of the soil. Denison [56] found that this equation is also applicable to the Bureau's data. This can be shown by plotting the logarithm of the average maximum pit depth on equivalent areas against the logarithm of the time of exposure. Within the error of the observations, the relationship is linear. This equation has the advantage over those of Fetherstonhaugh and Putnam that it need not be assumed that all pit depth-time curves have the same general shape.

In view of the general applicability of the Ewing equation, values of the constants k and n have been determined for the soils at all the National Bureau of Standards test sites for which sufficient data have been accumulated to give reliable values. The value of k appears to be a function of the fundamental corrosivity of the soil, and the value of n seems to be related to the degree of aeration. The degree of aeration of a soil largely determines the effectiveness of the corrosion products in protecting the metal against continued corrosion. Based on this relationship, values of n to be used in the absence of data from which they can be determined directly have been assigned [56] to classes of soil according to their aeration, as follows: Good, 0.188; fair, 0.345; poor, 0.473; very poor, 0.682. The use of three significant figures is usually not warranted by the precision with which the values can be determined for a given soil or class of soils. As a rule, probably not more than one significant figure should be used.

### V. RESULTS OF FIELD TESTS ON FERROUS MATERIALS

### 1. CAST IRON

## (a) AFTER 17 YEARS OF EXPOSURE

Table 29 shows the loss of weight, maximum penetration, and the weighted average maximum pit depth of specimens of 6-inch cast-iron pipe exposed for approximately 17 years to 19 of the less corrosive soils of the original National Bureau of Standards test sites. Care should be used in comparing the materials, as about a third of the delawaud specimens were buried 2 years later than the pit-cast specimens. As has been stated, the depth of the deepest pit is a function of the area from which it is chosen. It has been observed that for small areas consistent results are obtained by averaging a number of pits proportional to the area exposed. Part of the data have been weighted in this table by averaging

TABLE 29.—Loss of weight and maximum and weighted maximum penetration of 17-year-old east-won specimens.*

		Duration	I	Loss of weight	t.	Maxi	Maximum penetration	niion	Weighted r	Weighted maximum penetration ¹	netration ^b
No.	Soil type	of exposure	Northern pit cast, L r(1)	Southern pit cast, Z (1)	rn deLavaud, t, Z C (2)	Northern pit cast, L (1)	Southern pit cast, Z (1)	deLavaud, C (2)	Northern pit cast, L (1)	Southern pit east, (1)	deLavaud, C (2)
	Bell chay Dublin chay adobe Everett gravelly sandy loam Maddox silt loam Genesee silt loam	Years d17.58 d17.49 d17.46 16.94 16.92	02/ft² 6.01 10.12 0.90 5.16	92/ft ² 4.19 11.62 0.67 6.95 5.92	02/ft ² 4.72 12.79 0.64 5.94 5.64	Mals 152 119 22 106 112	Mils 82 154 30 86 110	Mils 104 90 56 115 87	Mils 110 114 19 88 88	Mils 72 132 27 27 78 103	Mils 78 68 53 88 88
	Hanford fine sandy loam. Houston black clay. Keyport loam. Merrimae gravelly sandy loam. Mismi clay loam.	d17.53 d17.58 16.97 17.19 16.99	4.12 8.99 9.93 0.47 2.46	3.41 6.45 8 92 0.81 2.51	1.23 12.00 9.12 0.73	94 226 175 27 132	70 148 67 73 112	92 54 86 68 92	85 168 96 22 98	62 130 62 38 97	.64 .65 .76 .01 .71
	Miami silt loam. Miller clay. Museatine silt loam. Rocrolk fine sand. Romona loam.	16.93 d17.63 17.04 d17.74 17.53	3.20 7.41 11.15 2.14 4.67	3.87 8.56 8.88 3.19 0.72	2.90 6.17 8.17 2.14	200 140 122 122	157 204 344 58 40 40	68 128 170 103	76 132 50 100	138 168 201 54 36	62 104 150 98
	Ruston sandy Joam Sassatrias gravelly sandy Joan. Summit silt Joan. Unidentified silt Joam.	d17.68 17.22 17.41 17.43	1.48 1.56 6.15 3.61	1.85 1.81 6.05 6.49	1.63 1.70 3.54 5.99	355 32 32 32	102 73 94 62	75 65 83 83	76 36 22 24 24	88 47 58	68 55 80 80 80

a See table 3 for identification of the specimens.

- Average of 4 pits on 1. Lo or Specimens, or 4 pits on each of 2 C specimens.

- The number in parentheses indicates the number of specimens removed from each site. When more than 1 specimen was removed the average value is given.

- The C specimens are approximately 15.5 years old.

- A parenge of 4 specimens.

- A parenge of 4 specimens.

- Data out 1 specimen only.

TABLE 30,—Loss of weight, maximum, weighted maximum and estimated average maximum penetration of 12-year-old cast-iron pipe.

[Data on 1 specimen for L and Z, and on two specimens for C]

	t, Z	$\sigma_m$	Mils	3.8	2.7	4.9	9.0	22.1	10.2	22.1	6.6	8 9	1.7	43 5	0.5	32.3	1.6	0.0 8.6 34.0
	Southern pit cast,	18	Mils	293.7	176.5	109.8	111.4	80.7	43.3	117.1	82.1 89.0	69 2	117.1	185.4	147.2	295.3	120.7	57 0 178.2 448.7
epth	South	40		0 ⊭1	0	# <u>1</u>	27 #		0	+2	07:	 ] #	(+2	0 6	77	0	#5	00
num pit d	st, L	g m	Mils	54.4	34.8	9.09	27.76	17.7	19.4	23.9	4.6	7.2	14.2	27.9	9.9	5.1	25.4	0.0 16.3 20.9
Estimated average maximum pit depth	Northern pit east,	18	Mils	158,4	295.5	60.0	2.711	139.9	225.9	106.9	46.6	69.2	137.9	78.4	124.9	205 8	9.701	100 9 162.3 339.9
ated aver	North	g		+5	7	T# ?	27	77	70	#	# #	0	#	0 č	7	1	H	# 0 0
Estime	C	p _m o	Mils	9.7	9.5	10.4	o.0	6.0	15.0	2.9	24.1 (f)	8.3	3.4	€2	8.1	3.9	37.7	3.2 5.4 13.8
	deLavaud,	i,	Mils	147.1 51.1	6.62	127.7	0.6/	63.4	195.8	43.0	84.0 (f)	70.3	54.0	(f) 95.1	8.16	69.4	119.4	52.3 111.2 246.9
	de	ąş		0#		2 5	Q H	#	<del>د</del> ه	+13	±2 (€)	0	9=	£#	7 7	0	11	900
unuu u	South-	cast,	Mils	224 87	147	06	OR 01	72	228	63	65	53	91	170	135	171	110	+0+
Weighted maximum penetration	North-	cast, L	Mils	123 57	120	90	10	87	217	89	43	50	110	148	92	187	81	156
Weigh Pe	deLa-	vaud,	Mils	115	45	108	10 61	83	156	43	42	ĭċ	39	74	69	61	06	98
tration	South-	cast,	Mils	301 93	176	103	10	1	239	462	89	09	114	189 69	147	259	121	440+
Maximum penetration	North-	cast, L	Mils	152 62	135	95	₽ <u></u> 2	149	240 74	75	66 59	99	131	154 56	103	202	102	
Maxin	deLa-	vaud,	Mils	127 48	01	126	2 6	8	179	20	46	29	47	: :#	85	69	26	106
ght	South-		oz/ft	13.05	4.18	10.08	10.47	8.01	30.81	5.29	2.27 4.52	6.15	6.87	9.13	4.84	4.56	10,19	PD.
Loss of weight	North-		oz/ft²	13.47	3.93	10.76	0.49	5.67	20.28	5.75	2.00	4.95	8.39	10.12 9.22	2.66	5.47	7.68	8.38
្ម	deLa-	vaud, C	oz/ft²	12.68	3.18	19.13	10.09	6.76	16.84	5.55	1.61	3.96	8.27	10.91	2.74	3.13	6.22	9.18
Dura-	tion of expo-	sare	Years	11.65	e12.10	12.00	*12.10 e19.00	11.52	11.76		$\frac{11.92}{12.10}$	11.76	e12.02	12.00	11.71	11.63	11.65	11.65
:	Soila No.			-2	အ	4,	o %	· -	ωģ	10	11	14	15	16	18	19	20	ដួននេះ

38.2 12.7 () 24.8	9.3 85.0 38.2	71.4	0 4 5	7.8	28.8	4.0	5.8
143.9	116.7 244.0 216.3	376.1	115.7	110.2	269.1	$\}$ 142.5	186.0
# 75°	# 5 # 1		<del>7</del> 7 :#	# 0 #1	0	# <del>1</del> <del>1</del>	+ 7
15.5 23.4 67.0 31.1	(f) 24.7 29.3	57.2	() 2.4		21.2	3.7	5.9
85.2 186 4 383.0 187.0	(f) 136.1 219.1	246.8	. (f) 	89.9	208.6	(f) } 65.7	132.9
# # # 1 # # # 2 12	£ 0	77	€ ∓	, 0 #1	0	(+1) (±1)	+2
3.9 8.2 9.0	(5) 8.3 15.2	5.5	©	3.9	13.2	41.0	6.0
45.7 102.7 65.1 (f)	(f) 84.1 145.6	72.0	(f) 107.5	67.0	137 3	223.8	87.8
± # €	E ## 3	9#	€ .0	# 15 # 15	+14	# - 1 0 # - 1 0	# €
90 157 1111 169 162	100 120 180 49	40	104	96 98	274	104	140 98 28
85 115 111 342 181	123 18 93 160 67	230	57 10 57	71	193	164	123 61 14
45 97 87 87	121 88 105 90	88 89	8 88	66	63	142	112 55 62
111 173 185 196 186	117 76 144 193 60	51	119 125	103	311	151	142 102 33
107 143 179 353 203	143 21 201 78	8 2 8	8 29	74	230	241	139 67 17
	136 52 74 120 100	888	106	49	76	178	128 68 72
3.32 4.94 8.17 38.43 20.83	9.81 3.76 6.45 19.84 7.51	6.53	20.33 14.39	13.55	30.39	3.97	23 75 8.08 4.27
3.47 4.40 6.65 36.05 31.40	12.52 2.34 4.00 16.18 6.97	1.90	9.28	11.40	17.79	3.27	6.24 5.55 3.47
3.66 3.52 10.82 5.85 53.26	9.28 4.40 4.24 15.85 7.59	3.62		18.56		3.76	12.58 4.16 4.10
11.67 11.52 12.02 9.60 12.04	11.62 *12.04 11.66 11.67 11.67	12.09	12.01	11.99	•12.03	12.02	11.73 12.00 12.08
88288	85883	135 36	88 88	40	42	44	146 146 47

a See table 19 for identification of soils.

b denotes the number of specimens on ontited in averaging because of unsatisfactory data. The plus or a minus sign before a value in the b column denotes whether the pitting of the omitted specimens was more or less than the average. A number preceded by a ± sign indicates the number of specimens for which there were no data.

of a denotes the standard error of 5.

The C specimens in this soil are approximately 10 years old.

That ansatisfactory for deformining the pit-depth—time curves.

Specimens lost; average pit depths at the close of 6 years.

D = Specimens destroyed; no weight-loss data.

four pits from each 6-inch specimen. The justification for this weighting will be shown later.

If it is assumed that the data are properly weighted, the data in the last three columns of table 29 may be compared with similar data for wrought pipe in table 36 to 37. Such comparison should not be made for specimens exposed for different periods even if reduced to time-rate bases. Likewise, it is not proper to compare unweighted pit depths for specimens of different sizes; for example, the average maximum-pit-depth data in table 30 with similar data for wrought pipe of the same age.

### (b) AFTER 12 YEARS OF EXPOSURE

Table 30 is mostly a rearrangement of parts of tables 4, 5, and 7 of Research Paper RP883, Soil-Corrosion Studies, 1934 [45], which reports the data on the examinations of all the 6-inch cast ferrous specimens buried in 1922.

The average maximum pit-depth data were derived in the following way. A procedure was adopted that involves a minimum of assumptions as to similarity of materials, areas, and rates of corrosion. The first step was to plot all of the pit depths for each material in each soil, and to draw arbitrarily smooth curves for the data representing the trend of the data with respect to the duration of the exposure. If no specimens were missing, 6 or 12 points, 1 for each of the L and Z specimens and 2 for the C specimens for each period of exposure were plotted. Points on this curve corresponding to the six periods of exposure were taken off, and the ratios of the last point to each of the others were computed. Then each plotted point or pit depth was multiplied by the ratio for the corresponding time of exposure, thus reducing ail pit depths to corresponding pit depth for 12 years of exposure. The average pit depth, the standard deviation, and standard error were then computed for these adjusted observations, that is, for each material in each soil.

The magnitudes of the average pit depth and of the standard error depend, of course, on the way in which the curve referred to was drawn, and it is probable that, in some cases at least, more representative curves could have been drawn. This would have resulted in smaller standard errors and more accurate average values. Nevertheless, the worst result of an improperly drawn curve is an inaccurate average and a large standard error. The two taken together are true for any curve, although they may not be sufficiently precise to be of value. As the purpose of the study was to determine whether or not the differences between the average rates of penetration could be accounted for by the dispersion of the data as indicated by the standard error, there was nothing to be gained by calculating the average rates of penetration for cases where the dispersion was obviously so great that a very large standard error would be obtained. In certain cases an average with a satisfactory standard error could be made by neglecting one or two points on the assumption that such points were the results of abnormal conditions with respect to soil or material. This, however, is a somewhat questionable procedure. Table 30 gives the average maximum pit depths for each cast-iron material in each soil for which a satisfactory standard error could be obtained by the method just outlined.

If all specimens were removed and data from them used, the number of observations or, in statistical terms, the size of the sample was 12. In some mildly corrosive soils two specimens were left in the ground so that they could be examined after a long period of exposure. A few

specimens were lost or destroyed, and a few pit measurements have been neglected for reasons stated above. The table includes a column showing the number of specimens that were omitted from the computations and the reason for the omission. A  $\pm$  sign followed by a number indicates the number of specimens for which no data were available; a + sign and a number indicate that one or more pit measurements were neglected because the values were so much greater than the estimated values for the period that an extremely large standard error would result if the data were included, and a - sign indicates that the pit depth was much less than the estimated value. It is possible, therefore, for anyone to revise the table by eliminating the soils from which data were deleted without completely recalculating the table.

It will be noted that the average maximum penetrations have not been computed for the cast specimens because there were so many soils in which the data were erratic that averages which would be satisfactory for comparisons could not be obtained. This unsatisfactory condition is due, in part, to the small number of specimens of the cast materials and, in part, to the great dispersion of the data. In some cases the dispersion is the result of extra-deep pits, whereas in others, the cause of the dispersion is the absence of pits of the expected depth. The fact that the cast-iron specimens were larger than the others may to some extent account for the greater dispersion of their data and for the greater maximum pit depths that sometimes appear. Table 33 may be used for comparing the materials listed in the table, but the unweighted data are not comparable without adjustment with similar data for specimens having different exposed areas. The pit-depth-area for relation will be discussed later. The standard error is an indication of the reproducibility of the data. Roughly speaking, if the tests were repeated, the odds against a second average differing from the first by more than twice the standard error are about 20 to 1.

The probability that two average values are really different may be determined from the equation

$$\sigma_D = \sqrt{\frac{{\sigma_1}^2}{N_1} + \frac{{\sigma_2}^2}{N_2}},$$

in which

 $\sigma_1 = \text{standard error of average No. 1.}$ 

 $\sigma_2 = \text{standard error of average No. 2.}$ 

 $\overline{N}_1 =$  number of observations in average No. 1.  $N_2 =$  number of observations in average No. 2.

 $\sigma_D = \text{standard error of difference.}$ 

If the difference between the two averages is more than  $2\sigma_D$ , the chances are about 20 to 1 that the difference is not due to chance. If the difference is more than  $3\sigma_D$ , the chances that the difference is real are about 997 in 1,000. To illustrate, for soil 1 the average of the deepest pits on 12 C specimens was 147.1 mils, with a standard error of 9.7 mils, whereas the average of the deepest pits on 6 specimens of southern cast iron, Z, was 293.7 mils, with a standard error of 15.2 mils. The standard error of the difference, 293.7—147.1, or 146.6 mils, is

$$\sqrt{\frac{(9.7)^2}{12} + \frac{(15.2)^2}{6}} = \sigma = 6.80, 3\sigma = 20.4.$$

It is, therefore, highly probable that the difference of 146.6 mils is significant. On the other hand the standard error of the difference between

the pit depths on the C and L specimens in soil 1 is 38.56, whereas the difference between the pit depths is only 11.3. The difference is, therefore, probably accidental. This is due to the large standard error of the L specimens. Statistical treatment of the Bureau's soil-corrosion data is not entirely satisfactory because of the dispersion of the data and the limited number of the specimens subjected to one set of conditions. and because the necessary assumptions as to the distribution of the data are not strictly correct. However, such treatment serves a very useful purpose in that it calls attention to the limitations of the data. Some idea of the magnitude of an investigation that would give entirely satisfactory data can be obtained from the fact that the Bureau has exposed over 36,000 specimens and from the magnitude of the standard errors of the data. Consideration of these facts will make it evident that to obtain satisfactory data, it would be necessary to increase the number of specimens subject to a single set of conditions, and to limit the varieties of specimens and conditions. Whether or not the same investment with these limitations would yield more useful information is debatable.

# (c) AFTER 9 YEARS OF EXPOSURE

The results of examinations of specimens buried in 1922, some of which appear in table 30, indicated that some soils were severely corrosive and that pipe materials more resistant to corrosion were desirable. In 1932 the Bureau invited manufacturers to submit specimens of materials believed to be suitable for service in 15 soils selected because of their corrosivity with respect to the commonly used ferrous pipe materials. The materials submitted are the bases for the so-called tests of corrosion-resistant materials. The tests will not be completed until after the war, but data so far obtained indicate in a general way what may be expected [57].

Table 31 shows the losses of weight and maximum penetration of six varieties of cast iron. The specimens were about 14 inches long and 1½ inches in diameter and were laid in a horizontal position across the trenches. The table is comparable only with tables 41 and 42, which show data on steel specimens of the same size and age. As table 31 is based on the performance of only two specimens of a kind, the data

cannot be treated statistically.

To provide a more reliable basis for comparing materials and soils, tables 32 and 33 have been prepared. For each of the four periods of exposure, the losses of weight and maximum pit depths for each material in each of 12 soils for which the data were complete were expressed as percentages of the averages of the loss of weight and the maximum pit depths in all of the 12 soils. The results for the four periods were then averaged to form tables 32 and 33. This permits a comparison of the materials in each soil and of the corrosiveness of each soil with respect to each material. By taking the value of N as 8, the number of specimens of one kind in each soil, the standard error of the difference for any two materials or soils can be computed and the significance of the difference determined. Without the computations it is almost obvious that some of the soils are more corrosive than others, and that the high alloy, E, is the only material definitely superior to the others.

Table 31.—Loss of weight and depth of maximum penetration of east-iron pipe exposed for 9 years. [Average for 2 specimens. Each ounce per square foot corresponds to an average penetration of 0.0017 inch.]

	Dura-	 Horizontally cast in sand mold	ly cast in	sand mold			Low a	Low alloy —			High	High alloy —
Soil type tool oogled, Rattled ex- p. p. posure Maximum Loss of penetration weight	coated, I. F. Maximum penetration	 Ratt Co. Weig	led, a of zht	Rattled, G,h Maximum penetration	I. Loss of weight	I. Maximum penetration	J. Loss of weight	J. Maximum penetration	C. Loss of weight	C. Maximum penetration	E. Loss of weight	E. Maximum penetration
Years Mils 9.47 nb75	Mils ab75	 2000		Mils b57	02/ft² 2.76	Mils 58		Mils 62		Mills n b66	02/ft ² 1.35	Mils 1533
Hagerstown noan	4250+(7) 250+(7) 44	 2008 2008 2009		250+(7) 250+(5,7) 52	37.34 4.05	74 240 74	22.10 27.10 12.92	215 233 49	29.70 21.88 2.37	23 216+ 250+ 24	0.72 14.62 9.91 0.66	181 1858 1863 186
Rifle peat.   9.24   a163 + 20.77     Susqueleared dity   9.53   73   7.08     Susqueleared dity   9.47   b00   8.10     Gridel marsh   9.55   131   6.14     Doese day   9.21   250 + D/341	*163 + 78 + 590   131   250 +	20.77 7.08 8.10 6.14 DC34		a175+ 78 ab81 114 250+	*18.15 7.54 6.61 *9.98	110 113 104 104 317+	0.89 0.89 6.37 a11.40	8130 80 85 85 85 246	13.07 7.21 6.92 2.56	*93 119 *80 b40	10.00 2.33 2.73 1.63	a45 a36 ab32 ab32
145 b118 250+(5,7)	145 b118 250+(5,7)	 2.56 D	~~~	131 b152(7) 250+(5, 7)	11.04 8.22 #01.41+	172 214 327 + (5)	+	156 b193 b266 +(5, 7)	14.01 10.74 45.74	161 b141 b195+(5,7)	* 12.55 * 13.20 52.33	b35 40 250+(5, 7)

• Data for individual specimens differed from each other by more than 50% of the larger value.
• Avenge maximum bit depths or loss of weight of the 1839 removals are greater. The shigh maximum value of the 1839 removals are greater.
• Avenge maximum pit depths or loss of weight of the 1839 removals are greater. The shigh maximum value of the 1941 removals are greater.
• Avenge maximum pit depths or loss of weight of the 1839 removals are greater. The shigh maximum value of the 1941 removals are greater.
• Avenge maximum pit depths or loss of weight of the 1839 removals are greater.
• Avenge maximum pit depths or loss of weight of the 1839 removals are greater.
• Avenge maximum pit depths or loss of weight.
• Dividences that both specimens were destroyed. The number in parentheses is the approximate loss of weight.
• Data for I specimen only.
• The loss of weight.
• Explain for I specimen only.
• The loss of weight with sand removed.

Table 32.—Average of the relative loss of weight of the cast materials for four periods of exposure.

[In percent]

	Rattle	ed, Ga	Low a	lloy, I	Low a	lloy, J	Low a	lloy, C	High a	lloy, E	
Soil No.	Rela- tive loss	Stand- ard error	Aver- age								
53	42	14	30	8	30	9	34	11	13	2	30
55	36	5	30	6	34	14	30	10	9	2	28
56	246	36	205	53	203	42	187	28	89	14	186
58	166	24	175	33	172	37	190	11	67	18	154
59 ^h	37	4	34	0	33	3	24	3	22	16	25
60	129	31	128	21	112	18	104	22	61	16	107
61	50	12	56	14	57	15	52	11	18	4	47
62	82	13	70	16	74	15	67	17	20	3	63
63	45	11	53	10	66	12	32	7	10	1	41
64	308	39	376	38	387	47	394	58	74	12	256
65	104	27	128	29	130	24	164	$\frac{26}{22}$	29	ც	111
66	74	16	73	7	99	6	118		42	5	81
Average	110		113		116		116		38		

^a See table 9 for composition of the materials.
^b Data for only 3 periods of exposure.

TABLE 33.—Average of the relative maximum penetration of the cast materials for four periods of exposure.

		Average	67 82 137 146 41	77 68 100 68 174	107 118	
	3, E	Standard	7 t 1 1 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	2200	F-10	
	High alloy, E	Relative maximum penetration	84 48 83 83 83 83	88 88 88 89 89 89	46 44	48
	loy, C	Standard	223 23 34 35	18 27 32	14	
	Low alloy, C	Relative maximum penetration	70 83 134 162 18	67 61 106 51 212	127 147	103
	lloy, J	Standard	14 29 29 3	23 23 23	98	
[In percent]	Low alloy, J	Relative maximum penetration	72 85 138 146 49	22 98 98 188	127 142	107
[In po	loy, I	Standard	11 11 35 26 6	35 35 35 35	23.3	
	Low alloy, I	Relative maximum penetration	57 79 145 156 55	86 91 119 75 208	126 132	111
	zd, G	Standard	9 38 48 11	24 113 29	8 14	
	Rattled, G	Relative maximum penetration	69 100 171 191 47	93 115 77 190	114 136	115
	ted, Fa	Standard	38 30 12 12	20 13 13 26	88	
	Sand-coated, Fa	Relative maximum penetration	84 100 177 156 52	89 105 74 198	101	109
	:	No.	53 56 58 59	61 62 63 64 64 64	66	Average

a See table 9 for the composition of the materials, b Data for only 3 periods of exposure,

## (d) AFTER 2 YEARS OF EXPOSURE

The cast-iron specimens exposed for 2 years consist of 12- by  $3\frac{1}{2}$ - by  $\frac{1}{2}$ -inch sections of 12-inch class 150 Super deLavaud pipe bolted to flat 12- by  $3\frac{1}{2}$ - by  $3\frac{1}{2}$ - by  $3\frac{1}{2}$ - inch charcoal cast-iron plates by means of steel and charcoal cast-iron bolts.

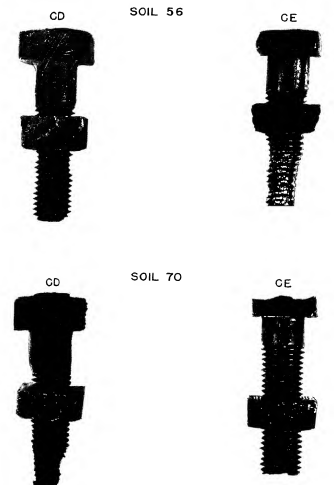


FIGURE 10.—Charcoal cast iron CD, and steel CE, bolts exposed to both soils 56 and 70 for approximately 2 years each.

Table 34 indicates that the charcoal cast iron lost more weight than the Super deLavaud iron in most soils, and in all but one soil the maximum pit depths on the Super deLavaud cast iron were definitely less

than on the charcoal cast iron. This might be attributed to galvanic action between the different metals in addition to the other causes of corrosion. Figure 10 shows the condition of the cast-iron and steel bolts connected to the couples in soils 56 and 70.

Table 34 .- Loss of weight and depth of maximum penetration of cast-iron plates and cast-iron and steel bolts (the plates were connected by the bolts).

			[Exposure	e 2 years]				,				
			deLa	vaud	Cha	reoal	Nuts	and bolts				
	Soil	Expo-		iron C		iron B	Steel CE	Charcoal cast iron,				
No.	Туре		Loss of weight	Maxi- mum pene- tration	Loss of weight	Maxi- mum pene- tration	Loss of weight	Loss of weight				
53 55 56 58 60	Cecil clay loam Hagerstown loam Lake Charles clay Muck Rifle peat	Years 1.91 2.03 1.91 1.91	oz/ft ² 1.21 1.03 6.13 5.98 4.27	Mils 30 32 45 42 23	$0z/ft^2$ 1.59 1.05 11.27 4.10 3.21	Mils 38 68 74 58 41	$0z/ft^2$ 1.92 1.28 20.10 6.47 5.06	0z/ft ² 1.81 1.56 9.04 7.12 3.32				
61 62 63 64 65	Sharkey clay. Susquehanna clay. Tidal marsh Docas clay. Chino silt loam.	1.92 1.90 1.88 1.90 1.91	1.66 2.52 1 94 4.53 1.89	36 34 36 36 34	2.56 2.73 2.69 4.64 3.39	58 57 57 57 54	2 54 2.57 (a) 3.28 4.24	2.28 3.31 (a) ^b 5.54 3.85				
66 67 69 70	Mohave fine gravelly loam. Cinders Houghton muck. Merced silt loam	1.86 1 90 1.90 1.90	$\begin{array}{c} 4 & 88 \\ 18.48 \\ 2.25 \\ 7.40 \end{array}$	34 88 38 96	6 28 20 67 2 08 10.50	54 102 36 122	16 22 1.71 11.19	(a) 18.18 1 86 10.12				

a Both specimens missing.

b Loss of weight for individual specimens different from each other by more than 50 percent.

#### (e) HIGH-SILICON CAST IRON

Included in the materials buried in 1922 were specimens of Duriron, a cast iron containing about 14 percent of silicon. The material is not machinable, but is used extensively for chemical plumbing. With a very few exceptions, the losses after from 10 to 12 years' exposure were negligible. There was no pitting or softening except along cracks in the specimens in soil 28, Montezuma clay adobe. Similar corrosion was found in this soil on two previous inspections, and on one inspection softening in spots occurred in Muck. Detailed data on high-silicon cast iron may be found in Research Paper RP883 [45].

#### (f) MALLEABLE CAST IRON

The corrosion of gray cast iron starts at and tends to follow the boundaries between the graphite plates and the ferrite crystals. The heat treatment which malleable iron receives slightly reduces the total carbon content and the free carbon takes the form of small rounded particles. The Bureau tests did not include a sufficient number of specimens to justify a generalization as to the relative corrodibility of grav and malleable cast iron.

Table 35 shows the data for the longest period of exposure. To the data on malleable iron have been added data on specimens of hightensile-strength cast iron and cast steel. Data on ordinary cast iron have been added for comparison. The malleable iron and cast steel specimens were in the form of elbows and with available apparatus it

## (d) AFTER 2 YEARS OF EXPOSURE

The cast-iron specimens exposed for 2 years consist of 12- by 3½- by ½-inch sections of 12-inch class 150 Super deLavaud pipe bolted to flat 12- by 3½- by ¾-inch charcoal cast-iron plates by means of steel and charcoal cast-iron bolts.

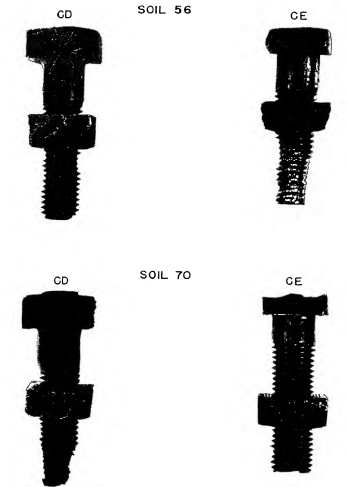


FIGURE 10.—Charcoal cast iron CD, and steel CE, bolts exposed to both soils 56 and 70 for approximately 2 years each.

Table 34 indicates that the charcoal cast iron lost more weight than the Super deLavaud iron in most soils, and in all but one soil the maximum pit depths on the Super deLavaud cast iron were definitely less

than on the charcoal cast iron. This might be attributed to galvanic action between the different metals in addition to the other causes of corrosion. Figure 10 shows the condition of the cast-iron and steel bolts connected to the couples in soils 56 and 70.

Table 34.—Loss of weight and depth of maximum penetration of cast-iron plates and cast-iron and steel bolts (the plates were connected by the bolts).

			[Exposure	2 years]								
			deLa	vaud	Cha	reoal	Nuts	and bolts				
	Soil	Expo-		iron C		iron B	Steel CE	Charcoal cast iron,				
No.	Type		Loss of weight	Maxi- mum pene- tration	Loss of weight	Maxi- mum pene- tration	Loss of weight	Loss of weight				
53 55 56 58 60	Cecil clay loam Hagerstown loam Lake Charles clay Muck Rifle peat	1.91	$0z/ft^2$ 1.21 1.03 6.13 5.98 4.27	Mils 30 32 45 42 23	oz/ft ² 1.59 1.05 11.27 4.10 3.21	Mils 38 68 74 58 41	oz/ft ² 1.92 1.28 20 10 6.47 5.06	$0z/ft^2$ 1.81 1.56 9.04 7.12 3.32				
61 62 63 64 65	Sharkey clay. Susquehanna clay. Tidal marsh Docas clay Chino silt loam.	1.92 1.90 1.88 1.90 1.91	1.66 2.52 1.94 4.53 1.89	36 34 36 36 34	2.56 2.73 2.69 4.64 3.39	58 57 57 57 54	2.54 2.57 (a) 3.28 4.24	2.28 3.31 (a) b5.54 3.85				
66 67 69 70	Mohave fine gravelly loam Cinders Houghton muck Morred silt loam	1.86 1.90 1.90 1.90	4.88 18.48 2.25 7.40	34 88 38 96	6.28 20.67 2.08 10.50	54 102 36 122	(a) 16.22 1.71 b11.19	(a) 18.18 1.86 10.12				

Both specimens missing.
 Loss of weight for individual specimens different from each other by more than 50 percent.

#### (e) HIGH-SILICON CAST IRON

Included in the materials buried in 1922 were specimens of Duriron, a cast iron containing about 14 percent of silicon. The material is not machinable, but is used extensively for chemical plumbing. With a very few exceptions, the losses after from 10 to 12 years' exposure were negligible. There was no pitting or softening except along cracks in the specimens in soil 28, Montezuma clay adobe. Similar corrosion was found in this soil on two previous inspections, and on one inspection softening in spots occurred in Muck. Detailed data on high-silicon cast iron may be found in Research Paper RP883 [45].

#### (f) MALLEABLE CAST IRON

The corrosion of gray cast iron starts at and tends to follow the boundaries between the graphite plates and the ferrite crystals. The heat treatment which malleable iron receives slightly reduces the total carbon content and the free carbon takes the form of small rounded particles. The Bureau tests did not include a sufficient number of specimens to justify a generalization as to the relative corrodibility of gray and malleable cast iron.

Table 35 shows the data for the longest period of exposure. To the data on malleable iron have been added data on specimens of high-tensile-strength cast iron and cast steel. Data on ordinary cast iron have been added for comparison. The malleable iron and cast steel specimens were in the form of elbows and with available apparatus it

was impracticable to determine pit depths on these specimens. Inspection indicated that the pitting did not differ greatly in form and extent from that on ordinary cast iron under the same conditions. A little additional information on the relative corrodibility of malleable cast iron and steel will be found in the section on the corrosion of bolts, table 50.

The specimens in table 35 differ considerably in the area exposed and in shape. The gray cast-iron specimens were much larger than the others, were buried at different times, and were exposed for somewhat different periods. The other materials seem to corrode somewhat less than the cast iron, but the differences are not large and may be accidental.

<i></i>		Dura- tion	Malle- able cast-	Cast	High-t	ensile- east iron,	Gray e	ast iron, L
Soil No.	Soil type	of expo- sure	iron ell, S. Loss of weight	E. Loss of weight	Loss of weight	Maxi- mum pene- tration	Loss of weight	Maxi- mum pene- tration
13	Hanford very fine sandy loam.	Years 10.16		$\frac{oz/ft^2}{19.81}$	$\frac{oz/ft^2}{10.87}$	$^{Mils}_{126}$	oz/ft²	Mil8
24 28 29	Merrimac gravelly sandy loam. Montezuma clay adobe Muck	10.63 9.60 10.08	13.44 10.58	4.02 16.03 11.69	1.06	100	4.74 *36.65 15.21	27 *353 145
42 43 45	Susquehanna clay Tidal marsh Unidentified alkali soil	10.05 10.73 10.55	$7.14 \\ 21.35 \\ 17.62$	7.44 8.69 19.94	5.73 $11.70$ $22.05$	104 163 161	11.27 11.42	172 157 232

Table 35.—Corrosion of malleable cast iron and cast steel.

## 2. WROUGHT FERROUS MATERIAL

This group of materials includes all rolled ferrous materials, such as open-hearth iron, wrought iron, Bessemer steel, and open-hearth steel, together with alloys, irons, and steels that have been rolled into their final forms.

# (a) AFTER 17 YEARS OF EXPOSURE

Table 36 and the weighted pit depths of table 37 are comparable with similar data in table 29 and present the data for the oldest wrought specimens. The weighting was based on the area of the specimens, that is, the single deepest pit on each 1½-inch specimen and the two deepest pits on each of two 3-inch specimens. As a check on this weighting, the corresponding pit depths of the 1½-inch and the 3-inch wroughtiron specimens may be compared. The same companies furnished both materials. Although the averages for all soils indicate that the weighting favors the 1½-inch material, there are 8 soils in which the weighted pit depths are greater for the smaller specimens compared with 10 soils in which the pit depths are less. The "e" and "y" specimens may be compared with the M specimens. It is possible that the amount of rolling or heating required to join the two sizes of pipe or the curvatures of the pipes may be a factor affecting the pit depths.

a Duration of exposure 11.67 years.

Table 36.—Loss of weight of 17-year-old wrought ferrous pipe.

foot,]
square
per
ounces
Ħ.
specimens
two
oţ
[Average

			-							
£,		Duration	1	1	1½-inch pipe			3-inc	3-inch pipe	A DESCRIPTION OF THE PROPERTY
No.	Soil type	of test	hearth iron, a	Wrought iron, b. d	Bessemer steel,	Bessemer steel,	Wrought iron, B. D.	Open- hearth steel,	Bessemer steel,	Open-hearth steel with 0.2% Cu,
97632	Bell clay, Dublin clay adobo Everett gravelly sandy loan. Maddox silt loan. Genesses silt loan.	Years 17.58 17.49 17.46 16.94 16.94	7.83 7.16 2.20 7.60 5.40	88.40 9.50 1.66 5.49	77.12 2.03 5.82 8.82	8.11 8.31 1.53 6.40	88.11 1.89 5.62	7.73		7.44 9.16 2.03 5.08
12 17 24 25	Hanford fine sandy loam. Houston black clay. Kaypot loan. Merrimac gravelly sandy loam. Miami clay loam.		7.18 7.94 7.94 3.02	12.39 8.78 1.88	9.54 10.95 1.79 2.70	0.35 10.41 1.40 1.40 2.00	12.51 12.51 1.30 1.30	5.39 15.98 9.40 1.31	5.10 11.21 9.51 1.56	5.20 7.57 7.24 7.24 7.44
323333	Miami silt loam. Miller clay. Muscatino silt loam. Norfolk fine sand. Ramona loam.		4.03 9.33 4.08 1.54	11.5.10 11.42 4.16 4.16 13.13	4.28 10.14 5.72 3.72	94.39 6.39 6.39 6.39	6.05 6.34 6.34 9.25 9.34	8.53.99 8.53.99 8.53.99	2 4.88.4.8 6.90.90.90.90.90	4 8 9 4 8 7 2 8 8 9 4 8 7 5 8 8 9 8 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9
36 38 41 47	Ruston sandy loam Sassafins gravely sandy loam Summit silt joam Unidentified silt loam	17.68 17.22 17.41 17.43	3.28 2.71 5.87 5.80	4.11 7.42 6.36	4.13 2.30 6.87 7.86	8.35 8.35 8.35	2.54 2.34 5.51 6.10	22.82 5.50 6.50 4.83	2.58 7.03 5.38	3.10 2.50 6.99 5.11

 $^{\rm a}$  Data for 1 specimen only.  $^{\rm b}$  Loss of weight of the individual specimens differed from the average by more than 50% of the larger loss.

TABLE 37.—Maximum penetration and weighted maximum penetration of 17-year-old wrought ferrous pipe,

[Average of 2 specimens; in mils.]

		#Mon"		Open-	0.2% Cu	steel, Y	64	8:8:2	95	29	323	###	49	6. 6.	47 56	53	31 70	#					
	Weighted maximum nonetesti	man benefit	3-inch pipe	Bessemer	steel,	M	56	4 19	22	83	778	25	50	823	31	48	82:	7.					
	iehted mavi		3-ii.	Open- hearth	iron,	М	62	12 12	53	¥8	948	54	57	#88	88	59	6198	Se .					
	We			Wrought	, mon	B, D	46 62	21.80	28	28	41	51	41	51	315	3.00	5.55 4.55 4.55 4.55 4.55 4.55 4.55 4.55	2					
				Open- hearth	o.2% Cu steel,	X	71	25	601	69 62	28 22	76	88	65 49	#	33.51	78						
-			3-inch pipe	Bessemer	, succi	M	82.5	77 48 89	70	52.55	38	96	83	92	37	49 35	101 48						
BIUI IIIIIII			3-mch	9-mc	%-W	- 9-m	Open- hearth	11.0II,	4	970 910 920	967	3 3	282	228	70	28;	388	£	288	80 <del>7</del>	_		
mode = va	Maximum penetration			Wrought iron,	R C		47 65 39	34.8	24	612	32.2	5 5	37.8	348	3 1	3%8	30 30	-					
.	Maximun					-			Bessemer steel,	>		42 20 20	19	64	36.55	26 43	2 %	374	183	11	985	57	
		1%-inch nine		Bessemer steel,	9		56 ⁵ 51 19	8.52	72	41	16	43	228	69	7.5	888	37						
		1½-in		Wrought iron,	b, d		9345	818	48	88 88 88 88	4 °C	43	84	34	50	88.35	53						
				Open- hearth iron,	cđ	i	72 27 27	969 	02	345	73	43	45°	120	51	122	42						
		Duration	of	cyposite		Years	17.58 17.49 17.46	16.94	17.53	16.97	16.99	16.93	17.04	17.53	17.68	17.22	11.40	a The meni-					
		מייו	No.			6	1002	o.	ខាដ	24.	25	38	385	35	36	814	;	a The meri					

a The maximum penetration and the weighted maximum penetration for the 112-inch pipe have the same value. b Data for 1 specimen only.

## (b) AFTER 12 YEARS OF EXPOSURE

Tables 38, 39, and 40 contain the same kind of information as table 33 and show the data for 12-year-old wrought pipe. Only the weighted pit depths on specimens of different size should be compared without adjustment. As a very rough average factor for comparing unweighted pit depths one may assume that for small changes in area, doubling the area increases the maximum pit depth on ferrous pipe about 10 percent. However, the pit-depth-area relation is somewhat different for different soils and materials.

At the bottom of the tables 39 and 40, average rates of corrosion for most of the soils are given. The difference in the soils is so great that average rates for all soils have little value, except that they permit a comparison of different materials exposed to the same conditions. These averages have been shown graphically in figure 11. As the pit depths were not adjusted to take account of the areas of the specimens, only materials of the same size are strictly comparable.

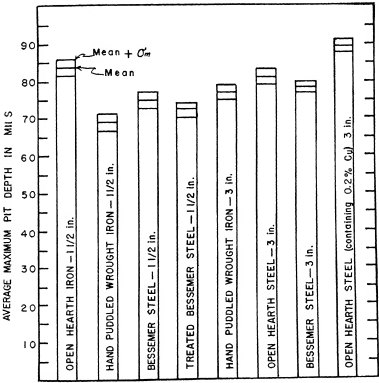


FIGURE 11.—Average maximum pit depth on wrought pipe for 12 years of exposure;
23 soils.

The figure 11 indicates that there may be a real difference between open-hearth iron specimens and the other  $1\frac{1}{2}$ -inch wrought specimens,

as the difference between the average pit depth of the iron and that of the other 1½-inch specimens is approximately twice the sum of the standard errors of the materials. Similarly, the 3-inch open-hearth steel containing 0.2 percent of copper may corrode at a different rate from that of the steel to which copper was not added. In both cases the reason for the differences may be the character of the surfaces of the specimens employed, and the conclusions might not apply to materials having similar compositions but different surface finishes. In the case of the open-hearth iron, the surfaces of the specimens were covered by an almost continuous thin oxide film which broke down in a relatively few places, thus concentrating the galvanic action. A basis for this suggestion is the low loss of weight of the specimens of this material.

The copper-bearing steel specimens, on the other hand, carried unusually heavy patches of mill scale at certain points on their surfaces. It is possible that galvanic action between this mill scale and the remainder of the surface of the pipe accelerated the corrosion, or that after a period of exposure the mill scale became loosened and galvanic action between the unprotected spots and the oxidized areas caused additional corrosion. Care should be exercised in generalizing from the

tests of the material from a single source.

The average penetration is smallest for the wrought-iron specimens, but the difference between the average maximum rates of penetration for wrought-iron and Bessemer steel is not sufficient to show positively a difference in the rates of corrosion of these materials for either the 1½-inch or the 3-inch specimens. Although it cannot be said that the materials do not differ, the figure indicates that on the average the rates of penetration do not differ greatly.

Figure 11 indicates also that the maximum pits are generally deeper on the 3-inch than on the 1½-inch specimens, but tables 39 and 40 show that this is not always the case.

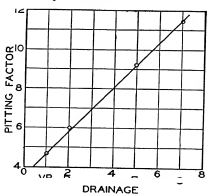


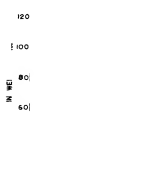
FIGURE 12.—Correlation between pitting factor and drainage.

One of the early observations concerning the soil-corrosion specimens was that the corrosion was not uniform and that this lack of uniformity was different for different soil conditions and different materials. The term "pitting factor" is used to designate the ratio of the maximum to the average depth of penetration by corrosion. The pitting factor is an

indication of the uniformity of corrosion and tends to become less as specimens grow older. For the specimens in tables 36 to 40 it ranges between 2 and 40. If the soils are divided into four groups with respect to drainage (aeration), as in table 5, and values 1, 2, 5, and 7 assigned to the terms "very poor," "poor," "fair," and "good," respectively, and if the pitting factors of all the wrought materials in each soil are computed from tables 32 to 40 and plotted against the values of the soil drainage, figure 12 results. The curve indicates that the pitting factor is lowest for the very poorly drained soils. The pitting factor gives no indication of the depth of the corrosion nor the number of deep pits, and it is not an adequate expression of the seriousness or the distribution of corrosion. It tends to be larger for larger specimens.

## (c) AFTER 9 YEARS OF EXPOSURE

Tables 41 and 42 show the loss of weight and maximum penetration of 9-year-old wrought specimens. These data are comparable with those in table 31 as approximately the same areas were exposed. Tables 43 and 44 were prepared in order to provide a better basis for comparison of materials by the same method as that used in the preparation of tables 32 and 33. The number of specimens of each material from each site was eight. The standard errors are so large that probably the difference between the materials is not significant. There appears to be a real difference in the corrosivities of the soils. Figure 13 shows graphically the averages given in tables 43 and 44, and indicates that with respect to the relative average values of all materials in the 10 soils for the 4 periods of exposure, nickel-copper steel (D) is better with respect to pitting and loss of weight. However, it should be noted that the mill scale had been removed from these specimens prior to burial. This might have had an effect on the pitting, as will be evident when the data on the nickel-copper steel given in the following section are compared with the data for a steel of similar composition, but from which the mill scale had been removed.



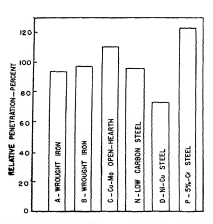


FIGURE 13.—Relative loss of weight and maximum penetration of wrought ferrous for four periods of exposure.

814786°--48---6

Table 38.—Loss of weight of 12-year-old wrought ferrous pipe.
[Average of two specimens, in onnoce nor source foot]

1.5

	1.1	verage or two	specimens, 1	A verage of two specimens, in ounces per square foot.	r square toot	7		-: 6		
		-		II-2-III	1½-incn pipe			3-inc	3-inch pipe	
	Soil type	Duration of exposure	Open- hearth iron, a	Wrought iron, b, d	Bessemer steel, e	Bessemer steel, y	Wrought iron, B, D	Open- hearth iron, K	Bessemer steel, M	Open-hearth steel with 0.2% Cu,
Allis silt loam. Bell clay. Ceci clay loam. Chester loam. Dublin clay adobe	m. dobe.	Years 11.65 12.01 12.10 12.00 12.10	8.92 85.95 3.71 7.03 6.46	8.77 7.12 7.12 4.90 6.61 7.58	7.03 85.81 4 69 6.70 7.03	9.29 5.92 5.14 6.23 5.46	9 56 6.32 4.62 7.11 6.42	9.45 5.37 4.17 6.96 7.31	a11.56 a6.59 4.34 6.92 7.32	8 81 5 91 7 7 4 7 7 1 4 7 1 4 7
Everett gravelly sandy Maddox silt loam Fargo clay loam Genesee silt loam Gloucester sandy loam	Everett gravelly sandy loam Maddox silt loam Targo elay loam Geneses silt loam. Gloucester sandy loam.	12.09 11.52 11.76 11.51 11.95	0.99 4.95 4.73 4.29	1.17 5.14 6 91 4.76 5.45	1.45 4 91 7.69 5.21 4.88	0.91 a4.30 6.48 4.96 4.35	1.12 4.82 8.66 4.89	1.07 4.91 7.94 5.03 4.38	1.52 5.64 8.29 4.58	. 1.3885;2 1.7885;2 1.2085;2
Hagerstown loam Hanford fine sandy Hempstead silt loan Houston black clay Kalmia fine sandy l	Hagerstown loam. Hanford fine sandy loam. Hempstead silt Joan. Houston black day. Kalmis fine sandy loam.	11.92 12.10 11.76 12.02	1.69 3.35 4.09 8.28	54.4.8 54.42 56.55 56.55 56.55	2.08 4.07 5.21 7.69 8.01	1.87 a3.92 4.54 7.83	1.97 4.01 *4.78 8.30 7.68	1.98 3.59 4.28 8.15 7.58	1 74 3.60 4.58 8.11 7.51	2.18 4.68 9.09 7.56
Keyport loam. Knox silt loam Lindley silt loam Mahoning silt loam Memphis silt loam	nm. agam. loam. sif loam. It loam.	11.78 11.71 11.63 11.65	9.45 2.95 2.90 6.63 6.97	8.61 2.67 3.50 6.54 7.64	8.34 3.82 6.10 7.52	9.02 2.64 3.42 6.01 7.06	10.56 2.03 3.25 5.63 7.93	10.22 2.73 3.45 6.12 6.89	9.65 2.40 3.32 7.78	9 96 3 88 3 48 7 65 7 00
Merced silt loam Merrimac gravelly s Miami clay loam Miami silt loam	Merced silt loam. Merrimae gavelly sandy loam. Misami elay loam. Misami silt loam. Miller elay.	12.09 11.95 11.67 11.52 12.02	20.14 1.26 3.40 3.53 7.08	21.13 1.36 *8.97 4.25 8.55	19.80 1.55 3.45 3.62 8.79	20.42 1.39 2.90 3.47 7.57	20.71 1.23 2.81 3.97 8.32	19.58 1.43 3.64 3.68 6.96	19 80 1.40 3 31 3.77 8.23	24.34 1.24 2.78 3.44 8.08
Montezuma clay adol Muck	Montezuma elay adobe. Muck. Musk. Muscatine sit loam. Norfolk fine sand. Ontario loam.	9.60 12.04 11.62 12.04 11.66	15.67 21.49 5.18 2.87 3.10	15.43 16.19 5.57 2.99 3.52	16.38 19.26 4.82 2.85 3.26	16.78 19.44 5.30 2.69 3.69	16.81 15.59 6.30 2.75 3.62	16.19 17.92 5.76 2.79 3.62	15.54 14.84 5.63 2.76 4.15	17.69 20.11 5.18 3.43 3.28
		_	_		_		_		_	

14.04	2.66	16.28
4.79	5.06	3.18
1.90	8.33	11.17
2.46	5.96	4.82
10.39	13.40	3.08
12.84 5.31 2.48 8.97	2.23 6.36 6.96 6.01 11.05	13.15 3.39 9.29 4.81 2.77
11.61	2.53	12.74
4.29	5.37	2.77
1.68	8.59	11.31
2.82	6.30	4.33
8.39	11.17	2.96
14.30	2.37	14.13
5.88	6.28	3.43
1.80	8.58	10.61
2.33	6.48	4.69
8.25	8.91	3.24
14.25	2.58	17.63
3.15	5.20	3.46
11.12	6.75	9.27
2.87	5.35	4.38
6.97	11.44	2.81
14.33	2.77	19.52
4 03	6.06	4.72
2.20	7.21	9.67
3.20	6.31	4.48
7.68	12.55	2.99
13.99	2.39	16.58
4.77	5.50	4.07
2.69	7.94	9.06
3.28	6.16	5.07
9.02	17.07	3.93
14.13	2.11	15.51
3.81	4.83	2.92
1.64	6.86	9.72
2.48	5.96	4.00
8.29	10.58	2.27
11.67 12.00 12.04 12.04 12.04	12.01 12.04 11.99 12.03	12.02 11.61 11.73 12.00 12.08
Peat. Pen silt loam Ramona loam Ransina sandy loam St. John's fine sand	Sussafras gravelly aandy loam Sassafras silt loam Sharkey eday Summit silt loam Susquehanna clay	43 Tidal Marsh. 44 Wabash silt Joans. 45 Unidentified alkeli soil. 46 Unidentified sandy loam. 47 Unidentified silt loam.
35 35 37	33 39 42 42	44 44 47 47

a Data for 1 specimen only.

TABLE 39.—Maximum, weighted maximum," and estimated average maximum penetration of 11½-inch wrought ferrous pipe exposed for 12 years.

		×	18	4.6	73.2 6.5	77.7 7.3	38.3 1.6		28 1 0 8 58.8 1.6	37.1 3.6		58.3 5 1 140 3 9 0	ο i ο	5.	59.9			67 4 3.0	23 1 1.8
			40	. 0:	# 0	# 67	+2		0 #	. 89	0	# 0	0 0	#	00	0	9#	0	# 2
	(2		-	8 9	3.4.	10	2 1		1001	. 88		12.2			4 C1		2.0	1.8	1.6
	acton (n = 1.	ð	41		* * * * * * * * * * * * * * * * * * * *	0.17	49.3	-	73.0	35.2		50 1 143 0		41.8	63.5.4	64.3	63.1	66.2	15.8
	na benefit		9	0 %	0	7 = 7	112		# 0	- 67 #	0	m 0 9	10	4	00	Ŧ	9=	0	<u>ا</u>
ll impagn	o maxima		0,11	5 9	- 67	0.9	5.		2.0 4 1	2.9	3.6	7.5	5.6	<del>-</del>	00:	5.1	2.8	1.4	1.0
ntod avora	anca average	p, d	1%	85.2	77.5	74.7	53.9		37.9 79.4	48.9	74.1	111.5	68.3	38.1	21.7 21.7 21.7	59.0	48.2	64.2	21.2
Retim	Estimated average maximum penetration $(n=12)$		9	06	107	7=	67		00	# 33	H ?	i O	0	#	00	0	9=	0	: - ::1
			o md	4.4		18 7	3.3		3.3	3.8	7:-	.00	9.0	6.6	ဝ က ဝ က	4.9	4.1	3 8	61.0
		et	:14	74.3 63.0	95.6	137.3	38 4		53.7 100.9	0.4	.0. .0. .0.	10901	84.0	41 0	63.2	57.4	7.17	65.0	25.3
			ęş	99	0	NI H	#		†°	67.67	0 0	000	0	#0	0	0	9#	0	4
g.		Treated Bessemer steel.	y	80	:G 6	3	37	18	81 82 ES	33	85 05 05	37.75	63	88	99	65		145+	123 8
enetratio	4	Besse- mer steel,	<b>.</b>	84 555	20 02	0	43	19	878	970	71	26	26	4=	I.E.	22		69 145+	25
Maximum penetration	1 11	Hand- puddled iron,	p'q	48	81	2	15	20	923	52	°75 °58	109 50	61	82	62	67		99 145+	£ 23
M		Open- hearth iron,	ಹೆ	93 95 96	N150+	-	88	23	100 43	20	99	101	83	33.4	žč	- 29		145+	858
٩	-kiri	tion of he test in		Years 11.65 12.01	12.10	3	12.10	12.09	11.76	11.95	11.92	11.76 12.02	12.00	11.78	11.63	11.65		11.65	11.95
	7	No.		-22	ω <del>-1</del>		o	16	~ & G	01		12	 9	17.	61	 8: ;	:	 읽器	# 150 # 150

4 8.00 8.00 8.00 8.00 8.00 8.00 8.00 8.0	2.3	9	4.9			3.4	3.0	2 1
48.5 146.9 121.5 55.5 40.8	49.5		71 6	54.4 86.8 85.4	95.2	78.5	19.8	72.4
++++	7 0 9	7	9 0	000 #	009	+2	# 2	
7.3 6.6 8.0 3.0	3.3	H	0.6	2.5	12.6	8.4	1.6	2.2
58.3 155.2 117.7 55.2 36.0	50.2		94.1	49.5 79.7 81.0	119.4 105.0 66.3	8.68	20.7	75 0
4 # # # # # # # # # # # # # # # # # # #	0 %		0	F000	000	7	# 5	:
2.4 7.88.7 1.2 2.0	3.4			6.8 6.7 12.9	7.8 9.3		3.9	2.3
147.8 99.8 50.2 33.9	50.5			49.7 86.7 112.4	95 2 86.9 56.4	51.0	27 0	69.1
+# # # +# 22 + 0 0 + + + 1 2 2 2 4 5 1 2 2 2 4 5 1 2 2 2 1 2 2 2 2 2 2 2 2 2 2 2 2 2 2	+10	#::#	0	F00	000 #	7	#3	
3.7 9.6 5.0	5.7		5.6	8.2 15.8 5.3	4.3 12.1 7.6	8.4	2.8	2.4
46.6 174.4 58.4 36.5	63.2	57.0	76.1	70.0 157.8 106.8	95.0 95.8 87.7	85.1	19.0	83.6
+ + : + + 2 - + : 2 - + : 2 - : 4 - : 2 - : 4 - : 2 - : 4 - : 2 - : 4 - : 2 - : 4 - : 2 - : 4 - : 2 - : 2 - : 4 - : 2 - : 2 - : 4 - : 2 - : 2 - : 4 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2 - : 2	° 7 0 1 0 1 0 1 0 1 0 1 0 1 0 1 0 1 0 1 0	11	0	00°	, # 000	<del>7</del> 7	#3	
60 145+ 145+ 51 31	50	32 °5 48	30	83 91	88 105 69	82 104	31	
61 137 + 136 + 59 34	46	41 18 49	79	20 20 20 20 20	111 100 63	85	26	
48 132+ 90 51 29	43	84 46 84 84	34	95 95	2888	78 62	46	
42 145+ 54 54 34	58	58 10 57	35	139	94 91 87	49	17	
12.02 9.60 12.04 11.62 12.04	11.66	12.00 12.09 12.04		11.99	12.03	11.73	12.08	
33 888 23	33 83	134 135 36	138 138	944	24.24	45	47 Aver-	ager.

The maximum penetration is the average of the deepest pits on each of 2 specimens. For 1½-inch pipe this has the same value as the weighted maximum penetration.
 Specimens of white of the average.
 A number in the 5 column preceded by a ≠ indicates the number of specimens for which there is no data,
 Z denotes the average.
 A number in the 5 column preceded by a ≠ indicates the number of specimens for which there is no data,
 Z denotes the average maximum pit depth.
 Data concess the standard error of Z.
 Data unsatisfactory for determining the pit-depth-time curves.
 Specimens lost; average pit depths at the close of 0 years.
 A plus (+) indicates that I or both specimens were punctured due to corrosion,
 A verage of the 28 soils, with no deletions because of erratic data.

TABLE 40.—Maximum, weighted maximum, and estimated average maximum penetration of 3-inch wrought ferrous pipe exposed for 1.2 years.

9 9 3 ထ က ထက်က 969999 ď ၀က မ დ 6.j 3.1 200000 10 17. ≃. 996 22770  $\succ$ :00 Ġ 9 9 0 7 8 10 104 75. 79. 85. 147. 81 48888 176 85.8 8 388 57 177777 **₩**0 054050 F779 40 Estimated average maximum penetration (n=12)1-010 4 9 1 6 4 C) 5.4 4: 0 2 0 4 4 . 00 10 വയുന്നുക က်လက် Ξ ~ oo 84.0 70.1 117.3 68.5 75.0 ∞∞ಣಣ≎ 18 Ħ :88 \$ \$ \$ \$ \$ \$ \$ 163 8233 8 0 1 8 0 900 4.3 6.7 10101401 ë. ဗ က းပဲ 222 20 000 1-010 - 10 01 10 10 167.1 M ដែ 113. 88 48 18 88.21. 86.22. 25.57 48884 63 4.4 3.23 0 mc 6.5 ٠. بر. ن e 0,00000 10. In mils] 00 to 44 00 to 994 0 B, D าให 23.85 89 ŧ2; 45. 86. 888 156. #38#8 3 7²7° 7 077770 # 0 0 0 0 0 0 0 0 0 ę, 2488 4 1 2 884 tr 528 322 Weighted maximum  $\succ$ 161 penetration 95 69 11 3 22 446 688 49 49 49 77 77 63 68 68 35.21 145 × 83 M 91 51 8 46 113 446 45 45 66 66 66 67 76 149 沒 А 146 **444** 7. \$ 33817 17 17 1888 9934 56 ď, Open-hearth steel, 152 49 63 55 55 Maximum penetration Besse-n mer steel, M 73 420 430 430 84 46 248148 81 65 72 72 70 \$499 65 163 Open-1 hearth steel, K 94 54 68 60 48 71 53 83 83 83 150 27178 Open-( hearth iron, B, D 28 28 28 28 83 57 58 58 58 44 84844 41 68 45 150 8 Years 11.65 12.01 12.10 8 2 95 31 95 95 10 200 8 Dura tion test ន 85 83 83 83 12 5 12. 12121 ==== Šći Š -- e1 e2 8~~@5 

2.8	4.6 8 9 5.1	10.8	8.3 6.4 6.0 6.0	13.1 13.1 2.2 2.0	1.8
36.7	$ \begin{array}{c} 76.3 \\ 221.1 \\ 61.3 \\ 40.7 \end{array} $	104.3	116.9 100.0 82.5 78.7	115.6 101.5 89.4 80.5 30.6	89.3
#2	# # # # # # # # # # # # # # # # # # #	#::# 0 #::#	00.00	7 3 3 5 0 0 H : + H	
2.9	12.9 5.1 13.1 2.9 2.5	2.4 18.9 	6.1 4.4 2.9	4.0 9.1 3.6 7.7	1.5
30.5	99.3 148.6 142.6 68.4 \$ 40.1	63 3 114.5	97.5 69.3 82.6 76.8	108.2 102.7 82.9 97.9	78.7
+ + 5	## # # 4 # + 2 # + 2	#5 #5	0 # 0 10 10 10 10 10 10 10 10 10 10 10 10 10	#+: +# 57:	
3.6	5.1 11.1 10.1 7.9	3.7 8.7 2.3	3.0 8.9 7.9 3.9		2.2
28.5	83.5 207.5 216.1 64.7	65.7 116.3	67.5 74.4 105.4 73.6	124.6 130.7 62.0 111.3	81.1
# # :	# # # # # # # # # # # # # # # # # # #	# # 5 # : # 7	00.00	## + # 0 0 (# + 1	
3.0	2.4.7 6.9 7.7 8.8	5.0 14.0	4 4 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8	25.6 6.6 5.3 5.3	2.0
24.8	58.9 169.3 126.4 60.2 43.1	62.6 131.3	72.1 64.2 70.2 86.8	92 2 138 4 63.8 86.2 86.2	76.8
# # :	# # # # # # # 5 5 7 7 8 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9	# 5 # 5 # 5	0 :002	++500	
30 45 45	65 216+ 171 63 36	90 55 43	120 22 98 74 77	106 68 71 77 24	
28 53	68 142 101 60 36	58 91 57 44	91 36 81 70 73	88 77 73 113 27	
3 24	76 180 194 51 39	81 98 35 19 59	60 22 69 87	124 106 55 110 57	
26 44 41	56 160 117 53 37	55 55 55 58 58 58	26 26 80 80 80	133 862 89 89 89 89	
31 52 43	74 216+ 4216+ 67 37	224883	127 26 106 75 80	116 72 75 124 80 80	
30 57 38	83 152 119 63 40	62 97 50 69 45	95 43 94 78 75	22 23 114 27	
20 46 46	81 1183+ 216+ 54 41	86 108 36 22 60	22 22 23 23 24 25 25 25 25 25 25 25 25 25 25 25 25 25	125 128 112 62 24	
28 44 44	55 167 134 53 41	59 104 55 37 ·	8280	34 735 238	
11.95 11.67 11.52	12.02 9.60 12.04 11.62 12.04	11.66 11.67 12.00 12.09 12.04	12.04 12.01 12.00 12.04 11.99	12.03 12.02 11.61 11.73 12.00	
24 25 26	30 28 31 32	22 24 25 25	41 41 41 41 41 41 41	224444 4	Aver-

a 5 denotes the number of specimens omitted in averaging because of unsatisfactory data. The sign of 6 denotes whether the pitting of the omitted specimens was more across than the average. A number in the 6 column preceded by a # sign indicates the number of specimens for which there were no data.
b \(\begin{align*}\) denotes the standard error of \(\beta\).
c \(\beta\) denotes the standard error of \(\beta\).
Data for 1 specimen only.
c \(\beta\) denotes the standard error of \(\beta\).
d \(\beta\) bata for 1 specimen only.
c \(\beta\) to determining the pit-depth-time curves.
1 \(\beta\) put (et ) indicates that 1 or both specimens of specimen only.
A \(\beta\) to determining the pit-depth-time curves.
1 \(\beta\) put (et ) indicates that 1 or both specimens were punctured due to corrosion.
4 \(\beta\) bata (et ) \(\beta\) bata (et ) in deletions because of cruite data.

Table 41.—Loss of weight of wrought pipe exposed for 9 years.

			21111111		In ounces per	square foots]					
				Vrought iron					Alloy steel		
	Soil		Mechani-	A -1	-В	Cu-Mo open-hearth	Low carbon	2.5% Ni	5% Cr,	18% Cr.	18% Cr 8% Ni.
No.	Туре	Hand puddled,	puddled,	Average	Standard error	iron,	steel, N	1.1% Cu, D	P	x	R
53 55 56 58 59 60 61 62 63 64 65 66	Cedi day loam. Hagerstown loam. Lake Charles clay. Carlisle muck. Rifle peat. Sharkey clay. Susquehanna clay. Tidal marsh Dosas clay. Chino all loam. Mohave fine gravelly loam.	7.80 *8.52 *16.00+	3.70 3.78 19.54 12.68 2.34 *16.48 45.66 9.38 4.24 f18.35+ 11.44 b9.99	3.71 3.73 21 18 12.62 2.39 15.37 6.04 8.59 6.38 17.18+ 12.52 7.90	0.3 .1 .5 .4 4.8 0.6 1.1 1.9		b4 00 3.82 28.76 16.24 4.70 16.72 5.78 6.65 9.03 D h12.86 18.56 58.39+	b2.53 b2.06 16.69 b6.55 b2.61 e10.28 b3.37 5.42 7.68 b e24.79 b6.00 e b7.54 D	b2, 24 1, 92 23, 58 13, 88 2, 76 15, 55 5, 10 4, 64 65, 85 D b10, 91 18, 32 27, 92	(d) 0.012 1.33 (d)	0.022

- a Each ounce per source for corresponds to an average penetration of 0.0015 inch. The maximum loss of weight for the 1939 removals is greater. The maximum loss of weight for the individual specimens of the 1939 removals is greater. The maximum loss of weight for the individual specimens of the 1939 removals is greater. Data cannot be used because of abnormal corrotion due to the presence of asphalt on the ends of the specimen.

  Data for the individual specimens differed from the average by more than 30%:
  Data for the perimens outlying the presenting was destroyed by corrosion.

  Data for 1 perimens outlying the yellow corrosion.

  A transpolose of weight of the 1939 removals is greater. The maximum loss of weight for the individual specimens of the 1941 removals is greater.

Table 42.—Depths of maximum penetration of wrought pipe exposed for  $\theta$  years.

	[In mils]												
	Soil Wrought iron							Alloy steel					
		Hand	Mechani- cally	A+	-В	Cu-Mo open-hearth	Low carbon	2 5% Ni	5% Cr,	18% Cr.	18% Cr 8% Ni,		
No.	Туре	puddled,	puddled, B	Average	Standard error		steel, N	1.1% Cu, D	P	x	R		
53 55 56 58 59	Cecil clay loam	*50 *60 *96 118 32	^{b73} ⁸⁴ ^{af} 106(7) ¹¹⁶ ²⁸	62 72 101 117 30	9 8 10 4	* d109 + 93 *145 +(7) *96(7) d22	59 59 154+(7) 110 440	*37 62 145+(7) *52(7)	70 *84 *136(5, 7) 111 20		8		
60 61 62 63 64	Rifle peat. Sharkey clay. Susquehanna clay. Tidal marsh. Docas clay.	455 61 472 100 5120+(7)	d64 d86 d101 d55 145+(7)	60 74 86 78 132+	16 16 22 19 12	461 82 486 b 467 145+(7)	d27 d96 d87 *54 154+(5, 7)	d38 a41 bd58 d70 e145+(7)	110 74 *70(7) 136 + 154 + (5, 7)	<6 *112 *96	<6 e #26 #36		
65 66 67	Chino silt loam	*102(7) * *88 145+(5, 7)	110 b130+(7) 145+(5, 7)	106 (7) 109 + 145 +	6	*106 145 +(5, 7) 145 +(5)	112 154+(5, 7) 154+(2, 5, 7)	74 ^{b d} 96 +(7) 145 +(5, 7)	b131+(7) 154+(7) 125+(5)	84	6		

- A verage pit depths of the 1939 removals are greater. The since naximum pit depth of the 1939 removals is greater.
  The since naximum pit depth of the 1939 removals is greater.
  The plus sign (+) in all cases indicates that one or both specimens were punctured.
  The plus sign (+) in all cases indicates that one or both specimens were punctured.
  The maximum pit for individual specimens differed from each other by more than 50 presents.
  The punctured is present to the presence of the present of the pit depth indicates that I or both specimens of a previous removal was punctured, i.e., (2) indicates a puncture after 2 years, etc.

  A number in parentheses after the pit depth indicates that I or both specimens of a previous removal was punctured, i.e., (2) indicates a puncture after 2 years, etc.

  Deeper pits present because of harmound corresion due to the presence of asphalt on the cale of the pips. These pits have not been counted against the pips.

Table 43 .- Average of the relative loss of weight of wrought specimens for four periods of exposure.

## [In percent]

		Wroug	ht iron		Cu-M	Cu-Mo iron,		Low-carbon steel,		steel,	5% C	r steel,	
0.11		A a	В		H		N N		D		P		A
Soil No.	Rela- tive loss	Stand- ard error	Rela- tive loss	Stand- ard error	Rela- tive loss	Stand- ard error	Rela- tive loss	Stand- ard error	Rela- tive loss	Stand- ard error	Rela- tive loss	Stand- ard error	Aver- age
53 55 56 58 59	62 56 206 153 31	14 9 37 19 2	63 56 179 153 28	14 10 24 22 1	60 45 162 158 33	8 6 31 25 1	62 51 256 176 48	6 8 47 27 3	43 33 133 118 35	8 6 21 18 5	35 26 192 132 34	4 3 46 27 4	54 44 188 148 35
60 61 62 63 65 Aver- age	132 78 86 74 167	21 15 3 11 26	137 73 98 56 146	21 14 9 8 13	120 61 86 69 207	19 11 9 4 14	175 62 92 102 192	20 14 10 8 18	$ \begin{array}{r} 117 \\ 45 \\ 60 \\ 59 \\ 97 \\ \hline 74 \end{array} $	22 4 8 10 18	142 55 56 60 182	39 11 7 5 19	137 62 80 70 165

a See table 7 for the composition of the materials.

Table 44.—Average of the relative maximum penetrations of wrought specimens for four periods of exposurc.

## [In percent]

		Wroug	ht iron		Cu-M	o iron,		arbon	Ni-Cu	steel,	5% C	r steel,	
		Aa	В		н		steel, N		D		P		
Soil No.	Relative maxi- mum pene- tra- tion	Stand- ard error	Relative maxi- mum pene- tra- tion	Stand- ard error	Relative maxi- mum pene- tra- tion	Stand- ard error	Relative maximum penetration	Stand- ard error	Relative maxi- mum pene- tra- tion	Stand- ard error	Relative maximum penetration	Stand- ard error	Average
53 55 56 58 59 ^b	96 108 107 112 37	11 13 15 18 5	103 115 117 117 29	9 15 17 23 3	134 130 128 123 17	9 15 28 30 6	86 93 138 132 43	9 10 31 29	56 76 130 103 14	2 5 37 23 5	100 117 207 117 37	11 4 26 9 10	96 106 138 117 30
60 61 62 63 65	62 65 106 84 150	6 7 12 17 8	66 68 134 59 158	6 17 23 5 9	54 85 144 101 185	13 24 23 14 29	52 84 126 68 126	18 20 18 12 8	49 53 94 61 98	12 8 11 12 7	110 60 140 144 186	11 14 19 11 8	66 69 124 86 150
Aver- age	93		97		110		95		73		122		

^a See table 7 for the composition of the materials. ^b Data for only 3 periods of exposure.

b Data for only 3 periods of exposure.

It should be pointed out that such graphs as figure 13, which are based on the averages of the performances of materials that do not behave the same in different soils, have limited value, as the material that shows the best average performance may not be the best for some specific soil condition.

# 3. FERROUS-ALLOY SHEETS AND PLATES

For convenience, many of the ferrous alloys submitted in the more recent tests were in the form of sheets or plates. Whether data on plates are applicable to pipes of the same material or not has not been determined. If the plates are laid flat in the trench as most of them were in the Bureau tests, the aeration of the under side may be poorer than that of the upper side. A large part of the specimens show maximum corrosion on the under side.

Table 45 shows the loss of weight and maximum penetration of seven high-chromium-alloy sheets after 9 years of exposure. With the exception of the 12- and 17-percent-chromium materials in three soils, the losses of weight and the pit depths are very small. Comparing the data for the plate K with the pipe R in table 42, which is similar in composition, the pipe lost weight more rapidly in all soils where comparisons can be made. However, the sheets K were placed on edge and were more highly finished. Specimens of the high-alloy sheet buried in soil 64 for 9 years are shown in figure 14. The behavior of these specimens is characteristic of what may be expected of these materials in poorly aerated, corrosive soils.

Materials S and T were included in the test to determine whether or not manganese could be used instead of nickel as an alloying element. The number of specimens is too limited to justify a conclusion regarding this.

It should be pointed out that the notations M and U appearing in table 45 and in some of the following tables may not represent real differences. It is often very difficult to determine whether a specimen is unaffected or whether metal attack has caused an increased roughening of the surface, because the surfaces were rough before burial. In any case, whether the pitting is listed as M or U, the amount of corrosion is negligible.

Tables 46 and 47 show the loss of weight and maximum penetration of alloy steel plates buried 4 years, and table 48 and figure 15 show the relative loss and penetration of the materials in terms of the loss of weight and penetration of open-hearth steel plates for 2 periods of exposure. The data are based on the average performance of 4 plates in each of 14 soils. There are objections to this treatment of data, since the value of the average might be unduly affected by an abnormally large pit depth and because one material might be superior in one soil and correspondingly inferior under some other soil condition. However, examination of tables 46 and 47 shows that these objections are not important with respect to the materials under consideration. Table 48 indicates that the addition of about 4 percent of chromium to an alloy reduces the loss of weight but increases the penetration. This is explained by the postulate that the chromium renders most but not all the surface of the metal more cathodic because of an oxide film which breaks down in a few places. This would result in a large cathodic and a very small anodic area and a high current density and rate of penetration at the small anodes. In many soils the corrosion products are not precipitated on the anode and corrosion is continuous.

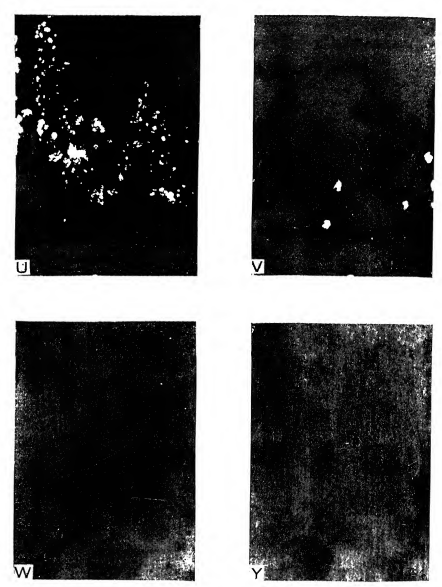


Figure 14.—Specimens of high-alloy sheet buried 9 years in Docas clay at Cholame. Calif. (soil 64).

U. 12-percent-chromium steel; V. 18-percent-chromium; W. 18-percent-chromium with 8-percent-nickel; I, 22-percent-chromium steel with nickel and manganese. The white spots on specimens W and Y are not holes.

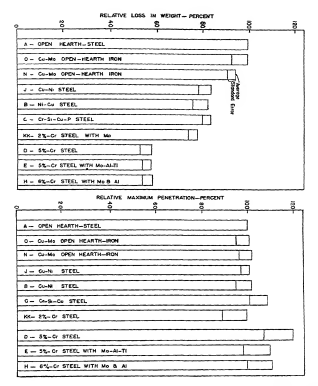


FIGURE 15.—Relative loss of weight and maximum penetration of ferrous plates for two periods of exposure.

d Average loss of weight or pit depth of 1939 removal is greater. 
• The number in parentheses after the pit depth indicates that at least 1 specimen of a previous removal was punctured, e.g., (2) indicates a puncture after 2 years. 
• Data for the individual specimens affilted from each other by more than 50 percent.

^a The number in parentheses indicates the number of specimens removed from each

test site.

^b Polished surface,

^c Data for 1 specimen only

Table 45.—Average loss of weight and maximum penetration of high-alloy steel sheets exposed for 9 years.

M, shallow metal attack, roughening of surfare, but no definite pitting.
If definite pitting, but no pits greater than 6 mils.
If a paperently wantered by corrowion.
If, apparently wantered by corrowion.
If, apparently an ance speciment ables because of corrowion, readering the companion of the exact penetration impossible. The thickness of the specimen bus been used as the maximum pit in this case.

1			oum ton, ton,		ls.	:		_		:	<u>8</u> +
	: (3)	12.94% Ni 1.50% Ni	Maximum pene- tration, average	. >-	Milk	-i-d	i ia	<u></u>		<u>:</u>	r P(2)
	7 1.0	121	Јања, аустике	:	02/Jts	0.0015		.0024		.0062	0020 1 12
	(3)	9.18% Ni 0.36% Mn	Maximum penc- tration, average	M	Mils	д	P	Ъ		4	P(2)
	) 69 %	2.0	Loss, аустадо	!	t)//20	0 00053	£2000	.0012		.0026	.00053
	(3)	8.95% Ni 0.44% Mn	Maximum pene- tration, average	Κħ	Mils	5	* 25 + (5, 7) TI M	M(5)	W	14+	^{d P(2, 7)} f 14 +(5, 7) P
	17.	ž ć	Lова, аvегада		£1//20	0.0010		.0012	.001	1.077	1.25 .0015
-	(1)	3.83% Ni 6.09% Mn	Maximum pene- tration, average	L	Mils					(T 4 P(2, 7)	M
	)	28	Ј.ови, а.устаде		z1f/20			:		0.44	.0042
and the second s	(3)	17.72% Cr 9.44% Mn	Maximum yene- tration, average	x	Mils					63+(2, 5, 7)	(2) M p
		7.71 9.4	Lови, в verage		1H/20			:		0.48	99000
- Andrews	(5)	0.09% Ni .30% Mn	Maxinum 1xme- tration, average	γ	Mils	n	Ъ	f P (5)		63+(2, 5,7,) 0.48	df 43+(2, 5, 7) 63+(7)
.	17.0	o	Loss, averago		z3f/20	0.0014		1.15		1.12	f 0.24 .70
the maximum pic in tons case.	(5)a 15% Cr	0.48% Ni .38% Mn	Maximum pene- tration, average	Ų	Mils	M	ď	dP (5, 7)		63+(2, 5, 7)	53+(2, 7) 55+(2, 5, 7)
IIIIXIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIII	11.9	0.4 E.	Ілж, вусгадо		oz/ft²	0.00042	.00084	d.0019		3.17	0.040
r arra		•	Soil No.		S	1818	59		623	22	65. 66.

TABLE 46,-Loss of weight of alloy iron and steel plates exposed for 4 years.

foot]
square
per
ounces
Ξ

	alloy steel, 18% Cr,	H	п0.48		
n steel	5.76% Cr 0.43% Mo 0.027% A1	H	1.44 1.00 14.32 n6.81 6.24	1.04 1.28 5.41 5.03 2.17	14.58 b12.88 0.72 10.25
4 to 6% chromium steel	4.67% Cr 0.51% Mo 0.030% Al 0.022% Ti	Ħ	1.59 1.19 16.26 7.16 6.23	1.81 1.09 5.15 5.11	12.04 be11.69 0.77 9.88
4 to 6	5.02% Cr,	Q	1.31 1.12 17.93 5.93 6.64	1.57 1.22 3.80 4.35 \$2.15	12.13 b18.09 0.86 b10.04
	2.01% Cr 0.57% Mo, 1	KK	2.18 2.38 14.48 8.36 6.00	4.67 2.99 5.39 4.53	12.26 °16.45 2.46 9.67
Low-alloy steel	Cr-Si- Cu-P steel 1.02% Cr,	Ö	2,41 2,33 20,76 9.03 10,23	4.59 3.05 4.14 5.12 5.44	13.56 °13.92 2.54 9.80
Low-all	1.01% Cu 1.96% Ni,	В	1.07 1.22 19.52 11.01 9.55	3.94 3.94 3.44 5.04	11.38 44.29 2.37 7.80
	0.95% Cu 0.52% Ni,	r	1.32 1.81 19.17 10.65 8.62	3.48 3.31 5.95 5.00	10.38 35.06 2.61 b7.62
rth iron	0.54% Cu 0.13% Mo,	z	3.02 2.73 17.37 9.36 8.22	5 65 4.69 6.63 5.03	11.96 27.70 2.90 b10.25
Open-hearth iron	0.45% Cu 0.07% Mo,	0	2.90 2.78 13.79 9.87 7.36	5.61 3.62 4.70 8.03 5.00	11.65 °33.22 2.86 b11.02
	Open- hearth steel,	Ą	3.23 2.58 18.38 9.88 9.52	5.40 3.71 n6.23 7.44 5.26	#16.82 34.27 4.21 10.63
	Expo- sure		Years 4.01 3.90 4.01 3.98	80.8 3.98 8.99	8.9888 8.9888
Soil		Type	Cecil clay loam. Hagenstown loam. Lake Charle clay. Muck Rifle peat.	Sharkey clay. Susquehanna clay Tidal marsh. Doeas clay. Chino silt loam.	Mohave fine gravelly loam Cinders Houghton muck Merced silt loam
		No.	55 55 50 50 50 50	64 64 65 64 65	96 67 70

Duta for I specimen only.

• Duta for I specimen only.

• Duta for the birdividual specimens differs from each other by more than 50 percent.

• Average loss of weight of the 1939 removals is greater.

Table 47.—Maxemum penetration of alloy iron and steet plates exposed for 1

	Hiph-	alloy steri	18% Cr,		F					
	1 Hford		6.76% Cr 0.43% Mo 0.027% TS	11.	158	197	2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	822	56 117 bf gdd	83
	4 to 6% chromium steed	4.67% Cr	0.51% M ₀ 0.030% Al 0.022% Ti.	E	50	9.7g	98	65. 66. 66.	, 88 1, 52	ьь 106
r I years.	4 10 5		5.02% (¹r,	G	57	95 b 344 b51	36 52	205	169 1657	121
ene secre pares exposed for 1 years.		3	0.57% Mo,	KK	223	642 622 622	35 56	45 ²	130+ b e68	161
b Martes	Low-alloy steel		1.02% Cr.	ī.	325	P22 P07	44	422	b80 b47	F6q
	lla-wo.l	1.01% Cu	1.96% Ni,	ä	1222	25. 28.	69 8	288 288 28	bf #84	78
In mile]		0.05% Cu	0.52% Ni,	£	50 96 96	b64 0	888	3%8	185 b f #90 15	82
,1	will iron	0.45% Cu 0.54% Cu	0.10% Mio,	2 2	100	n26	54 49 47	76 *57	198 b e74 16	271
_	Open-hearth iron	0.45% Cu	, , , , ,		116	82	988 148 148 188	7.5 6.5	e f #55 20 20	;
		Open- hearth	steel,	92	b100 b61		26 26 26	510	abe132+ 222 b77	
		Soil	Type	Cecil clay loam	Lake Charles clay. Muck.	Sharkey clay	Susquehanna clay Tidal marsh Docas clay			
i.			No.	533	8888	3 2	288	99	286	a Dat

b Dista for the individual specimens differed from each other by more than 50 percent.

b Dista for vorsion—impossible to measure true penetration.

d Amark in lad cases indicates I or more specimens punctured.

That corrison at I and.

Bad corrosion at I and.

Bad corrosion at I and.

Bate entirely destroyed at I end.

Bate for I specimen—impossible to obtain data from other specimen because of extreme uniform corrosion.

Average pit depths of the 1939 removals are greater.

Table 48.—Relative loss of weight and maximum penetration of alloy iron and steel plates.

[Average of two periods of exposure, in percent]

		Los	s of weight		Maximum penetration		
Symbol	Material	Aver-	Stand- ard Stand- devi- ard ation error	Aver- age	Stand- ard devi- ation	Stand- ard error	
A O N J B	Open-hearth steel Open-hearth iron—0.45% Cu; 0.07% Mo. Open-hearth iron—0.54% Cu; 0.13% Mo. Low alloy—0.95% Cu; 0.52% Ni. Low alloy—1.01% Cu; 1.95% Ni.	100 100 95 84 83	38 19 25 36	100 101 102 101 102	31 35 24 36		
KK D E	Low alloy—Cr-Si-Cu-P steel—1.02% Cr Low alloy—2.01% Cr; 0.57% Mo	84 78 58	19 20 27	109 100 120	45 58 70	11 13	
н	0.030% Al; 0.022% Ti	58 59	24 26	110 111	66 61	12 11	

The behavior of the 4- to 6-percent-chromium steel in this test is in agreement with the behavior of the 5-percent chromium-steel specimens in the 9-year test, where lower loss of weight and deeper pitting was observed in comparison to plain steel.

Further inspection of figure 15 shows that the nickel-copper steel (B) is no better than some of the other low-alloy steels with respect to loss of weight and pitting. This is contrary to the results of the 9-year specimens, where the nickel-copper specimens of approximately the same composition appeared to be somewhat better than the other specimens. It should be noted that the scale on the 9-year nickel-copper steel specimens was removed, whereas the 4-year specimens were coated with a hard, uniform layer of mill scale. The acceleration of pitting resulting from local failure of the oxide coating would account for the relatively poor performance of the 4-year nickel-copper specimen (B), as compared with the 9-year specimen (D) of the same composition.

Figures 16 and 17 show the low-alloy plates exposed to soil 70. Areas where the mill scale was left intact on the nickel-copper (B) and the

copper-nickel (J) steels are clearly shown.

Table 49 shows the averages of the loss of weight and maximum penetrations for a low-alloy plate and pipe and two high-alloy plates buried approximately 2-years. Although the loss of weight and depth of maximum penetration of the two high-alloy materials is slight, there is a consistent difference favoring the higher alloy. It is too soon to reach definite conclusions as to the relative merits of the copper-molybdenum steel and the chromium-molybdenum steel. The reader is cautioned against comparing these data with data for other materials exposed for the same length of time but buried at an earlier or later date, as two periods equal in length may differ considerably in amount or distribution of rainfall. For longer periods of exposure these differences will be of less importance.

## 4. BOLTS

Users of mechanical joints for pipes have frequently found severe corrosion of the bolts used, especially at the threaded ends. This led to the inclusion of a variety of bolts in the tests, some of which connected pieces of cast iron when buried, whereas others were completely exposed

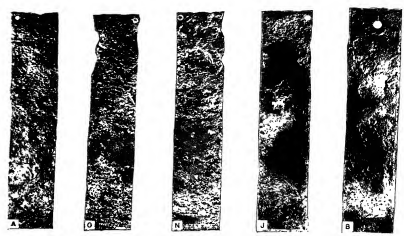


FIGURE 16.—Ferrous plates exposed 4 years to Merced silt loam at Buttonwillow, Calif.

See table 7 for the composition of the specimens.

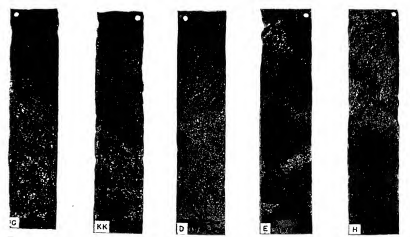


FIGURE 7.—Ferrous plates exposed 4 years to Merced silt loam at Buttonwillow, Calif.

See table 7 for the composition of the specimens.

to soil. None were under severe stress. The irregular shapes of the bolts, their partial protection by the nuts and the materials which they held together, the galvanic couples involved, and the relatively few specimens of each material subjected to the same conditions resulted in quite unsatisfactory data, some of which however, are presented in table 50. The pit-depth measurements were limited to the heads of the bolts because it was impracticable to measure the pits on the body of the bolt ac-

curately. The data indicate little difference in the performance of the different varieties of bolts. Sherardized bolts, and to a less extent leadcoated bolts were more corrosion resistant than corresponding steel bolts. which, incidentally, were submitted as wrought-iron bolts.

Table 49 .- Loss of weight and depth of maximum penetration of low-allow and high-alloy steel specimens exposed for 2 years.

U, apparently unaffected by corrosion. M, shallow metal attack, roughening of the surface but no definite pitting. P, definite pitting but no pits on either specimen greater than 6 mils.

Soil		Expo-	Mo o	u, .07% open- h iron .te), M		.49%		teel te)	Ni (pl	Cr, 11% steel ate),
No.	Type		Loss of weight	Maxi- mum pene- tration	Loss of weight		Loss of weight		Loss of weight	
53 55 56 58 60	Cecil clay loam	Years 1.91 2.03 1.91 1.91	oz/ft ² ^a 2.32 0.92 11.16 4.70 3.44	Mils 42 44 60 48 16	oz/ft ² 1.97 0.92 13.81 5.88 4.08	Mils 65 59 78 46 18	$0z/ft^2$ $0.0048$ $.0065$ $.0033$ $.0017$	Ü	0z/ft ² 0.0054 .0061 .0067 .0053 .0038	Mils M M M P P
61 62 63 64 65	Sharkey clay. Susquehanna clay. Tidal marsh Docas clay. Chino silt loam.	1.92 1.90 1.88 1.90 1.91	2.32 2.96 2.08 4.09 *3.14	41 40 22 47 43	1.98 2.57 1.76 3.92 3.87	42 40 38 40 51	.0032 .0020 .0046 .0018 .0048	ממממ	.0063 .0074 b.0084 .0063 .0067	P
66 67 69 70	Mohave fine gravelly loam. Cinders Houghton muck Merced silt loam	1.86 1.90 1.90 1.90	6.64 25.97 * 2.25 10.07	71 128 * 13 92	9.02 17.75 1.89 12.81	°145+ 124+ 22 130+	.0041 .0042 .0018 .0035	ממממ	.0095 .0056 .0060 .0078	P U U M

Data for 1 specimen only.
 Data for 3 specimens.
 +Indicates that 1 or both specimens contained holes due to corrosion.

## 5. EFFECT OF CORROSION ON BURSTING STRENGTH

The users of low-pressure pipe have frequently found on uncovering a pipe to repair one leak that punctures existed in several other places, some of which did not leak until the corrosion products were removed. If soil conditions are such that the corrosion products are deposited on the corroding areas, these products may form plugs of sufficient strength to prevent leaks until corrosion has weakened a considerable part of the pipe wall. This is shown by the following tests. A set of corrosion specimens, which included four pieces of 6-inch cast iron, were buried in Imperial clay near Niland, Calif. (soil 113). When the specimens were removed after an exposure of about 6 years, two of the cast-iron specimens were so corroded that they broke while being cleaned. The maximum pit depths in the other two were 0.243 and 0.345 inch, respectively. At the beginning of the test, a 36-inch length of 6-inch cast-iron pipe with the ends closed was filled with water and buried at the end of the trench. A small copper tube was brought out from the end of the pipe, by means of which hydraulic pressures have been applied to the pipe at 2-year intervals. Fifteen years after the pipe was buried it withstood a pressure of 400 pounds per square inch. A similar test has been conducted in Merced silt loam and in a muck soil in New Orleans similar to the soil at site 29 and in Houston black clay (soil 15) with similar results. In 1940, similar tests on standard and light weight

Table 50.—Corrosion of %-inch botts.

			Mal	leable-iron b	Malleable-iron bolts (4 inches long)	ng)		į		į	á -
Soil	Soil type	Number of	Duration	Decarb	Decarburized, Aa	Not decar	Not decarburized, Ba	High str	High strength, Ca	Steel D	Steel bolts, D
5		of each kind	of test	Loss of weight	Maximum penetration	Loss of weight	Maximum penetration	Lose of weight	Maximum penetration	Loss of weight	Maximum penetration
15	Hararatoum loam	6	Years 0 11	Grams 5 96	Mils	Grams	Mils	Grams	Mils	Grams	Mils
356	Carlisle muck	101	9.12	7.46	88	5.89	12	6.57	282	98.9	12
19	Sharkey clay	63	9.53	9.95	51	11.08	43	5.25	29	5.18	202
79	Susquehanna clay	7	9.47	8.05	88	8.93	22	9 74	28	8.10	72
	Average			7.68	48	7.75	43	7.35	47	6.21	33

Coated steel bolts and nuts (3-inches long)

Total loss of weight	Lead-coated steel, Steel, Db	t Bolt Nut Bolt Nut	us Grams Grams Grams Grams G.50 2.02 24 99 8 4.8	1 43 0.78 3.01	18.93 7.92 23 63	6.52 2 62 15.30	13.05 7.54 37.32	42.23 14.40 16.71	6 14.78 5.88 .11.47 7.04
	Sherardized steel,	Bolt Nut	Grams Grams 3.20 1.23						11.76 4.76
	Duration of test			10.63	10.08	10.05	10.73	10.55	
Number of	Number of specimens of each kind			₩-	41		41 .	4	
	Soil type		Hanford very fine sandy loam	Merrimae gravelly sandy loam	Muck	Susquehanna clay.	Tidal marsh	Unidentified alkali soil	Average.
	Soil No.								

a On head of bolt only.

b Included for comparison,

steel pipe were started. The light weight pipe blew out in 1943 at three sites at pressures of 151 pounds or less, but the standard-weight pipes withstood a pressure of 400 pounds per square inch. These tests indicate that corrosion products tend to plug the small-diameter pits and thus to render the pipes serviceable in spite of severe local corrosion. However, too much reliance should not be placed on the strength of corrosion products. If large areas are corroded deeply, or if the pipes are subject to severe water hammer or vibration, breaks or leaks may develop.

A somewhat different but related study of the effects of pitting has been made by a manufacturer of wrought pipe [58]. A number of lengths of steel pipe used underground and pitted by corrosion were cleaned. Heads were welded on the ends and hydraulic pressure applied until the pipes burst. The report concludes that the results of these tests indicate by a comparison of the actual bursting strength with the calculated theoretical bursting strength, that although the reduction in bursting strength of any given size and wall thickness depends upon the extent to which corrosion has occurred and the depth of the pits, it is significant that a single or a dozen or more widely scattered pits of varying depths up to one-half or more of the original wall thickness does not impair the bursting strength and usefulness of the pipe to any marked degree, as the surrounding wall supports the base of the pit and prevents the pipe from bursting, except at high pressures, that is, pressures many times the normal working pressure of the average water line.

It will be noted that the report [58] deals with the supporting strength of adjacent metal and unpunctured pipe from which the products of corrosion have been removed. No doubt, the tests first reported in this section were affected by the same factor, but it is probable that at some of the test sites corrosion has punctured the pipe wall, as specimens which have been similarly exposed, removed, and cleaned have been punctured after much shorter periods of exposure.

# VI. INTERPRETATION OF RESULTS ON FERROUS MATERIALS

#### 1. GENERAL CONSIDERATIONS

In interpreting the data obtained in the National Bureau of Standards soil-corrosion investigation, the reader should keep in mind the fact that the original purpose of the investigation was to determine the extent to which soils are corrosive with respect to metals commonly placed in them, in the absence of electric currents from street railways or other sources of power. It should also be understood that many of the phenomena related to the behavior of metals underground were not generally understood at the time the tests were undertaken. The results of the tests have been entirely satisfactory as far as the original limited objectives are concerned. Furthermore, certain more extensive inferences can be drawn from the data, provided always that the specific conditions under which the tests were carried out are given due consideration. Care should be taken, however, to guard against adopting too far-reaching conclusions from too limited data. It should also be emphasized that predictions based upon the behavior of individual test specimens cannot be relied upon with certainty except for similar materials operating under identical conditions. The performance of two specimens of the same material exposed to nominally the same conditions may differ considerably owing to unknown differences in material or environment. For this reason, it should be fairly obvious that general conclusions should be based on the average performance of a considerable number of specimens exposed to the same soils under as nearly identical conditions as possible.

Several factors that affect the corrosion of metal underground have already been mentioned, including depth of burial, scale on the surface, diameter and area of pipe, and duration of exposure. To these should be added the roughness and degree of oxidation of the surface, the condition of the soil at the time of burial, and other factors that may be

peculiar to a given location.

The data on loss of weight and pit depths shown in the foregoing tables give a fairly clear picture of the corrosion of the specimens used in the tests, and give a good indication of the extent to which corrosion occurs in the various soils. Also, by employing one or more of the empirical formulas already discussed, the progress of corrosion can be expressed. The accuracy to which a given equation represents the actual course of the corrosion process depends upon the dispersion of the data as indicated by the standard errors, which in turn depend upon the number of observations and their accuracy. If it is desired to make predictions of the performance of a pipe line buried in a given soil, it will be necessary to employ some method of extrapolation. It should be understood that extensive extrapolation is a somewhat hazardous procedure at best, especially with respect to corrosion phenomena, as it involves not only the extension of areas and time of exposure, the effects of which can be approximately expressed mathematically, but also of changes in environment which cannot be so expressed. However, as the environmental changes tend to counteract each other, statistical treatment of the data results in a fairly reliable determination of what may be expected on the average. What will happen in a single exposure is practically unpredictable. The standard error of the predicted result is a measure of the reliability of the prediction and should always be considered along with the prediction.

## 2. EXTRAPOLATION OF DATA

Any of the equations for the relation between pit depth and time may be combined with any of the equations representing the relation between pit depth and area. The choice of which equations should be combined depends somewhat on the ease with which the resulting equation can be applied to the data at hand and on the precision desired for the calculations. Although the use of different equations will yield different results, in many cases the standard errors of the differences resulting from the application of different equations to a set of data indicate that the differences in the results may not be significant. When the extrapolations by means of different equations are small, the differences in the results are often not very important. When the extrapolations are large, the standard errors of the results are large, and large apparent differences are therefore to be expected. The equations should be regarded as means for roughly estimating the corrosivity of soils and for this purpose are very useful. In every case the standard error of the estimate should be computed in order to give an indication of its reliability.

For the purpose of illustration, Ewing's pit-depth-time equation has been combined with Scott's and with Ewing's pit-depth-area equations. This gives the equations

 $P=kT^nA^a$  and  $P=kT^n$  ( $C \log A+1$ ),

in which P is the pit depth on an area, A, at any time, T, and C, a, k, and n are constants derived from an analysis of the data on pit depths. Table 51 gives the values of these constants for 47 soils. By means of the equations given above, the average depth of the deepest pits on 1,000 linear feet of 8-inch pipe exposed 30 years were calculated. By rearrangement of the equations, it was possible to calculate the average length of pipe associated with a puncture in 30 years and the time required for an average of one puncture of the pipe wall per 1,000 feet of pipe. These values are also shown in table 51. As the data for pure open-hearth iron, wrought iron, Bessemer steel, open-hearth steel, and open-hearth steel containing 0.2 percent of copper were not definitely different, they were averaged for the calculation of the constants in table 51. The data for cast iron were more erratic and were not used on this account.

The values of  $k_{5.3}$ , column 5, are the averages of maximum pit depth on specimens having areas of 0.4 square foot. Their exposure was 5.3 years. The values of  $k_{5.3}$  range between 21 and 107 mils. The values of the area factor, a, column 3, range from 0.08 to 0.32. The values of the time factor, n, column 7, range between 0 and 0.92.

Attention is called to the values of k, a, and n in order that the reader may realize how greatly soils differ, and also to the standard errors of these values so that he can see the erratic nature of soil corrosion, even under conditions that are somewhat less diverse than those frequently encountered by pipe lines.

In columns 11 and 15 of table 51 are shown the computed averages of the maximum pit depth to be expected on a group of 8-inch pipes 1,000 feet long and 30 years old. The standard errors in columns 12 and 16 are a measure of the variations that may be expected from the listed pit depths. The standard errors  $(\sigma_p)$  for the pit depths derived by Scott's equation are computed by means of the equation

$$\left(\frac{\sigma_p}{P}\right)^2 = (2.3\sigma_a \log A)^2 + (2.3\sigma^n \log T)^2 + \left(\frac{\sigma_k}{k}\right)^2$$

The standard errors for the pit depth derived by Ewing's equation were calculated from the equation

$$\frac{\sigma_p}{P} = \left(\frac{\sigma_o \log A}{C \log A + 1}\right)^2 + (2.3\sigma_n \log T)^2 + \left(\frac{\sigma_k}{k}\right)^2.$$

In these equations  $\sigma$  is the standard error of the quantity indicated by the subscript. The other letters have the same significance as in the other equations. In table 51 the unit of area is 0.4 square foot and the unit of time is 5.3 years. The large values of the standard errors of P show that columns 11 and 15 indicate only the order of magnitude of the pit depths to be expected under the assumed conditions.

Columns 13 and 17 show the computed average length per puncture for a 30-year-old 8-inch pipe line having a wall thickness of 0.322 inch. Columns 14 and 18 show the estimated time required for the development of an average of one leak per 1,000 feet on 8-inch pipes having a wall thickness of 0.322 inch. It can be shown that if the extrapolation of the pit-depth data is large with respect to either the area exposed or the period of exposure, the standard error of the estimate is correspondingly large. As many of the computed lengths and times are very long, their standard errors are very great, and consequently they indicate only that the corresponding soils are not very corrosive.

TABES 51.—Calculated data on pit depths and leaks on a pipe line.
Length of unit section 1,000 feet, diameter of pipo 8,025 inches, thickness of pipo wall 322 mils

1	uation	Time for puncture per 1,000 feet of pipe	18	Years 25 130 190 22 22 74	00 160 55 1,700 62	51 6,500 74 10 190	64 · 9,400 4,000 140 63	24 1,200 8 8.8 X10 ⁸ 780
	Ewing's pit-depth-area equation	Length of pipe per puncture in 30	17	Feet 3.1×10° 1.0×10° 1.9×10° 9.1×10 1.8×10°	2.4×10 ⁴ 1.4×10 ⁴ 1.4×10 ⁸ 1.0×10 ⁵	1 4×104 2.4×109 8.8×103 4.0	7.8×104 8.7×1015 8.2×1012 2.0×108 1.2×105	1.7×10 ² 6.9×10 ¹³ 0.7 1 1×10 ²⁰ 7.7×10 ⁴⁵
1	's pit-de	Stand- ard error	16	Mils 53 42 42 36 130 61	22 24 54 64	208 110 39	63 26 35 17 58	52 111 144 194 16
	Ewing		15	Mils 351 195 235 390 211	266 266 169 237	272 159 275 545 178	252 108 134 188 234	364 149 640 80 125
mils]	Juation	Time for puncture Deepper 1,000 est pit feet of in 30 pine years	Ξ	Frars 13 60 31 11 11	97 97 120 20 20	20 670 4 5 110	31 2,900 1,300 83 22	15 650 4 2.1×10 ⁴ 480
Length of unit section 1,000 feet, diameter of pipe 8,025 inches, thickness of pipe wall 322 mils)	Scott's pit-depth-area equation	I sugth of pipe per puncture in 30 years	13	Feet 6.2×10 6.1×10³ 1.0×10³ 2.7×10 2.0×10³	2.4×10 ⁷ 1.0×10 ³ 4.9×10 ³ 3.7×10 ³ 4.2×10 ³	3.9×10 ² 1.7×10 ⁴ 1.6×10 ² 4.0 6.4×10 ⁴	1.0×103 1.0×106 4.4×106 2.8×104 4.7×102	4.5×10 2.7×10 0.8 2.9×10 9.2×10
s of pi	ıpit-de	Stand- ard error	12	Mils 103 76 73 208 102	20 30 51 97 127	78 284 181 241 155	88 48 47 28 120	81 16 334 42 19
thickne	Scott	Deep- est pit in 30 years	11	Mils 475 254 320 572 290	75 188 353 97 378	367 215 465 770 212	319 136 164 226 371	480 169 863 121 144
inches,		ØC.4	10	0.069 .099 .082 .068	.026 .026 .032 .099	.071 .109 .055 .123	.053 .053 .026	.033 .024 .140 .128
ipe 8,025		ريم	6	0.417 .370 .413 .459	.409 .309 .365 .511	.396 .398 .573 .472	.320 .360 .301 .270	. 351 . 212 . 391 . 515
eter of pipe		$\sigma_m$	∞	0.06 .08 .05 .17	.05 .07 .15 .13	95:23:05	E. 25 25 25 25 25 25 25 25 25 25 25 25 25	70:03 60:09 90:09
set, diam	ants	*	1	0.49 .34 .17 .59	0 .46 .32 .16 .42	8. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1.	25. 11. 12. 13. 13. 13. 13. 13. 13. 13. 13. 13. 13	961212195
n 1,000 feet, d	Constants	okins	9	22.22.20 7.22.40	1.1 2.0 0.9 5.4 3.2	7.2 1.4.0 7.8 7.8 8.5 8.5	3.7 1.9 5.8 1.6 3.0	3.1 1.1 2.9 1.6 2.0
it section		kus	ũ	58.5 45.4 68.8 51.6 37.0	21.7 31.1 64.4 44.0 37.2	62.9 51.2 63.7 85.5 51.4	66.0 33.1 46.0 51.9 34.7	55.7 107.3 22.7 40.8
gth of un		g-#	4	0.021 .030 .024 .020	.009 .009 .028 .027	.031 .033 .033 .015	.017 .038 .018 .010	.012 .009 .044 .035
[Len	ļ	a	က	0.146 .131 .144 .160	144 116 133 172 184	140 1158 1158	118 111 111 103 188	. 129 . 082 . 139 . 172 . 088
To the state of th	Soil	Type	Ø	Allis silt loam. Bell clay. Geoil clay loam. Chester loam. Dublin clay adobe.	Everett gravelly sandy loam Maddox sift loam Fargo clay loam Genesee sift loam Gloucester sandy loam	Eggerstown loam. Hanford fine sandy loam. Hanford very fine sandy loam. Hempstead silt loam. Houston black clay.	Kalmia fine sandy loam. Keyport loam. Knox silt loam. Lindley Clay loam. Mahoning silt loam.	Marshall silt loam. Memphis silt loam. Merced silt loam. Merrimae gravelly sandy loam Mismi clay loam.
Ш	l	No.	1	H0100410	109878	122243	16 17 18 19 20	222222

85 27 10 7	84 13 39 120	5.2×10 ⁴ 28 3.3×10 ³ 35 23	90 48 17 17 16	26 550
3.6×10° 4.4×10° 0.02 .50 6 ×10°	9.1×10 12 2.2×10 5 2.6 7.9×10 3	1.2×1017 7.8×102 5.6×1014 3.2×103 1.8×103	1.3×107 1.5×104 2.8 2.6×104 3.6	7.1×102 1.2×1010
93 76 362 154 100	22 51 197 96 96	11 60 15 45 76	78 14 90 52 174	28
211 343 865 783 187	117 229 585 280 229	114 328 109 297 365	210 279 605 260 530	331 134
49 115 8 3 3 55	8 28 19 28 29	1.5×104 4 750 18 12	61 27 22 22 11	24.2
5.5×10 8 6.6 0.01 0.6 1.3×10 4	3.0×104 8.6×102 2.7 2.3×102 1.2×102	8.6×10° 6.4×10 1.2×10° 1.7×10° 4.5×10°	1.7×104 7.3×102 2.8 5.3×102	6.2×10 2.1×10
125 179 437 455 139	49 120 338 144 381	20 121 38 78 124	95 26 274 116 200	96 84
262 511 1,059 1,211 233	179 330 868 406 638	134 544 154 419 503	245 332 1,013 353 716	540 270
.091 .130 .041 .152	.095 .138 .109 .042	.041 .063 .095 .051	.055 .026 .081 .140	.040
.336 .511 .277 .538 .351	.508 .502 .500 .447	.255 .570 .443 .424	.252 .256 .590 .440	.762
2822	86. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1.	.00.00 .00.00 .00.00		86.
4.88.89.83	0 33 45. 55.	12. 23. 55. 56.	.30 .30 .78 .78	81.05 81.05
7.1 13.2 2.0 7.0	22.0 7.1 1.3	0.60 0.7.4.4.0.	8.04.0 8.4.04.8	4.4
45.7 38.1 86.0 92.0 32.1	40 4 44.8 56.4 41.0 26.5	45.4 65.4 27.5 47.4 60.9	54 8 84.7 83.2 58.5	79.3
.028 .037 .043	.027 .039 .028 .012	.014 .017 .028 .015	.018 .023 .038 .025	.030
120 170 106 178 123	.172 .165 .168 .157 .318	.097 .191 .153 .150	.095 .098 .195 .148	.186
Miami silt loam. Miller clay. Montexuma clay adobe. Muok. Muok.	Norfolk. Ontario loam. Peat. Pen silt loam. Ramona loam.	Ruston sandy loam. St. Johns fine sand. Sassfras gravelly sandy loam. Sassafras silt loam. Sharkey clay.	Summit silt loam. Susquehanna, clay. Tidal marsh. Wahash silt loam. Unidentified alkali soll.	Unidentified sandy loam
332328	32,33,33	38 38 40 40	44444	47

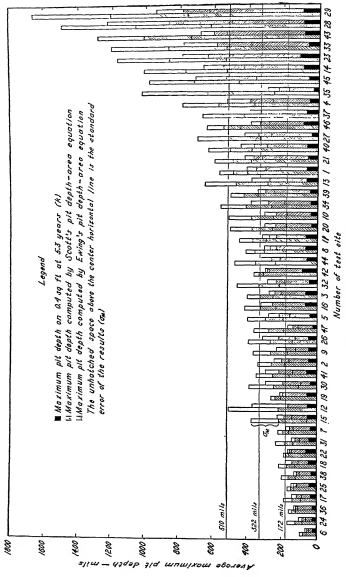


Figure 18.—Estimated maximum pit depth on 1,000 feet of 30-year-old 8-inch pipe.

Figure 18 which is plotted from data in table 51, shows graphically the estimated maximum depth of pit on 1,000 feet of 30-year-old 8-inch pipe and the standard errors of the estimates. The standard errors of the predictions are such that in most cases the values predicted by one equation fall within the averages predicted by the other, plus or minus once or twice its standard error. For smaller extrapolations, the results obtained by the two equations would of course differ less.

Although the data in table 51 must be recognized as rough approximations, they are of use to those interested in selecting the proper thickness for a pipe wall or in determining whether or not a protective coating

should be applied.

The relative depth of pits on pipes in different soils varies with the area of the exposed material and with the period of exposure. If the data in column 5 are compared with those in columns 14 and 18, it will be seen that although after 5 years of exposure the specimens in Cecil clay loam, soil 3, were pitted more deeply than those in Chester loam, soil 4, a pipe in the first-named soil would last much longer than one in the other soil.

One hundred seventy-two mils has been suggested [59] as the permissible thickness for 8-inch water-supply pipe for pressures up to 250 pounds per square inch. An examination of column 11 indicates that within 30 years such an unprotected pipe would average at least one puncture per 1,000 feet in 38 of the 47 soils. According to column 15 punctures on the same length would occur in 34 of the 47 soils. A good protective coating would of course reduce the number of punctures. As supply lines frequently are readily accessible, the cost of repairing them may not be great, and it is quite possible that the cost of repairs would be less than the cost of a protective coating or of thicker pipe. The table therefore does not indicate that light-weight pipe would be unsatisfactory. Column 15 indicates that standard-weight steel pipe (0.322 in.) would develop a puncture in 14 soils and class B cast-iron pipe (0.51 in.) in 7 soils under the same conditions.

It already has been said that Ewing's pit-depth-area equation probably represents available data somewhat better than Scott's. Table 51 shows that Ewing's equation results in the prediction of shallower pits,

fewer leaks, and longer pipe life.

The standard errors for the estimates of pit depths on 1,000 feet of pipe at 30 years are so large, however, they indicate that the estimated depths are quite uncertain. It probably would be desirable to recompute the estimates on some basis involving less extrapolation. One engineer limits his extrapolation to 40 feet of pipe.

## 3. LIFE OF PIPE

The expression "life of pipe" should be avoided because it has no generally accepted meaning, although the term is frequently used. By making a sufficient number of repairs, a pipe may be made to last indefinitely, although it might be more economical to replace it. Figures 19, 20, and 21 illustrate different types of repairs to steel pipe lines. Usually a pipe-line operator repairs or replaces badly corroded sections of his lines before leaks occur and so adjusts his reconditioning program that the number of leaks on his lines does not increase with the age of the line. Under such a policy, the life of a pipe line is the time there is need for the service.

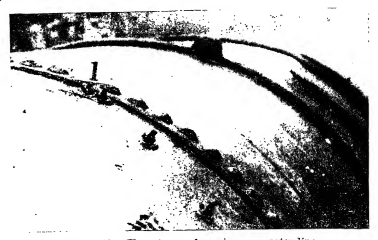


FIGURE 19.—Three types of repairs on a water line.

Bolts and washers used to close holes in large diameter steel pipe. Note also corrosion of rivet heads and bands used to hold patches.

Obsolescence rather than corrosion usually determines the time a pipe line remains in service, and the time required for a section of pipe to be punctured usually does not determine the time the pipe section will remain in service. This is a function not only of the depth of the deepest pit but of the number and depth of the other pits and the relative cost of repairs and replacement.

If by "life of pipe" is meant the average time required for corrosion to puncture a pipe wall, the answer is that it depends on the soil, the thickness of the pipe wall, and the area exposed, and the life of ferrous pipe can be estimated by means of the equations presented in the preceding pages. The life of a single section of pipe like that of an individual person cannot be predicted with any degree of assurance.

## 4. COMPARISON OF TEST DATA WITH DATA ON PERFORMANCE

Much has been said about the value of experience in estimating the corrosivity of soils. It is difficult, however, to express experience in sufficiently quantitative terms to make the information usable in the way that the results of tests are used, because the conditions under which the data of experience are obtained are often poorly defined, and because observations of identical conditions are usually not numerous enough to permit an estimate of the standard error or reproducibility of the experience. When experiences are analyzed, it is often difficult to determine their exact significance or their applicability to some different condition. Experiences are frequently contradictory and may be the results of some unrecognized conditions. They should, therefore, be examined carefully if they are to be applied to new construction.

As the National Bureau of Standards soil-corrosion tests were conducted with the cooperation of operators of pipe lines who furnished the test sites and local labor, it might be assumed that a comparison of the Bureau's data with the field experience of the cooperators would furnish a key to the application of the results of the test to pipe-line corrosion. When such a comparison is attempted, however, a number of

difficulties are encountered. Although there are pipes in the vicinities of most of the test sites, their extent in the soils of the test sites can be determined only by extensive soil surveys. Most of the pipe lines originally carried some kind of protective coating. In many cases this coating was very thin and imperfect, although in some instances the coatings were quite heavy. In certain cases the lines have been reconditioned and the kind of coating changed from time to time. Most pipe lines carry

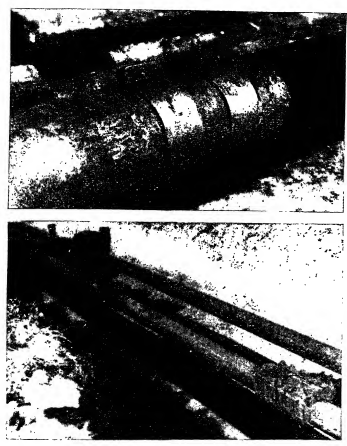


FIGURE 20.—Two types of leak clamps.
Top, home made, and bottom, commercial leak clamps.

currents which tend to protect certain portions of the line at the expense of other portions. Pipe networks in cities are frequently affected by stray currents from street railways, which modify the corrosive effects of the soils. The connection of one variety of pipe material to another, such as the connection of copper, brass, lead, or galvanized steel to cast iron, wrought iron, or steel mains, introduces additional galvanic cor-

rosion. The passage of a line through two different soils sets up a galvanic current. The pipe that lies in the same type of soil as the test specimens may be buried deeper or shallower, and thus lie in a soil horizon that is quite different in texture, aeration, acidity, or resistivity from that of the specimens. The interconnection of a new pipe with an old one sets up a condition that may accelerate the corrosion of the newer line. Two pipes of the same material in the same street or right-



FIGURE 21 .- "Half-sole" (welded patch) method of pipe repair.

of-way may corrode at different rates. In many cases records of the pipe lines are incomplete. In other cases the owners are reluctant to disclose their experiences because of possible effects on the valuation of their lines or the rates they may be allowed to charge for amortization as common carriers. For these reasons and others, no general comparison of the test data with the conditions of pipe lines has been attempted. However, a few comparisons will be made where conditions are especially favorable for such comparisons, although in no case are the conditions affecting the pipe line identical with those affecting the test specimens, and it is impossible to evaluate the effect of the differences accurately.

Soil 1. Allis silt loam.—Severe corrosion of service pipes in this soil has been reported, but definite data are not available. Ewing's equations indicate an average of one leak per 1,000 feet of 8-inch steel pipe

in 25 years.

Soil 8. Fargo clay loam.—This soil is similar to much of that in the city of Winnipeg, where severe corrosion of cast-iron water mains has been reported. Originally, the corrosion was attributed to stray-current electrolysis, but the extent of the corrosion attributable to stray currents has not been established. Ewing's equations indicate that in this soil an 8-inch steel pipe should develop in 30 years an average maximum pit depth of 266 mils on a 1,000-foot length.

Soil 9. Genesee silt loam.—A 6-inch steel line paralleling the test site was renewed because of corrosion after 14 years. Ewing's equations

indicate that this soil is not corrosive.

Soil 15. Houston black clay.—Severe corrosion of cast iron and steel has been reported in this soil. Table 51 indicates that this soil is not very corrosive.

Soil 23. Merced silt loam.—Leaks developed on an 8-inch steel line in this soil within 5 years. Table 51 shows this soil to be very corrosive.

Soil 27. Miller clay.—Frequent leaks developed on a 12-inch line in this soil in about 9 years. The data in table 28 are from observations on such a line in Miller clay somewhat better drained than that in which the specimens were buried. Ewing's equations indicate that an 8-inch steel pipe should average one leak per 1,000 feet in 27 years.

Soil 28. Montezuma clay adobe.—This is said to be a very corrosive soil, but no specific data have been obtained. Table 51 shows this to be

one of the most corrosive soils tested.

Soil 29. Muck.—Water mains in the vicinity of the test site corrode rapidly. The corrosion was attributed to electrolysis at one time, but the actual cause of the corrosion has not been definitely determined. Table 51 shows this to be one of the most corrosive soils tested.

Soil 32. Ontario loam.—The test site lies within a few feet of a 38-inch steel water main having a 56-inch wall. Within 2,000 feet of the test site this line developed 25 leaks in 42 years. The line was protected by a japan varnish baked on. The length of the pipe in this soil at this location is 7,480 feet, in which length there were 50 leaks, or an average of one leak for 150 feet in 42 years. According to Scott's pit-depth-area equation, the National Bureau of Standards data indicate there should have been an average of one leak in 5,349 feet of pipe. The formula of Ewing would indicate that the length per leak should be greater. The Bureau's data show that in this case the soil is much less corrosive than experience proved it to be. However, this steel main is paralleled by a 36-inch wrought-iron main having a 36-inch wall which has developed no leaks in this soil in 61 years. The difference in the two experiences is greater than the difference between the test data and experience.

The difference in the performance of the two pipe lines may be explained in several ways. The materials used were different, as were the protective coatings. It has been shown [60], in certain cases at least, that when a new pipe is connected to an old one, the latter is protected at the expense of the former. As for the difference between the results calculated from the tests and the leak records of the two pipe lines, it is obviously impossible for the experimental results to agree with both experiences. Most of the leaks on the steel line occurred on the top three-fifths of the pipe. This is rather unusual, as most cases of severe corrosion occur on the bottom of the line. The severe corrosion of the steel line may be the result of a difference in soil conditions at the top and the bottom of the pipe or because the coating on the top of the pipe was injured by the backfill. The record of these pipe lines illustrates very well how conditions not duplicated in a test may alter the results when the material tested is used in a practical way.

Soil 35. Ramona loam.—The specimens at this site are laid adjacent to a 6-inch east-iron gas main that has carried gas for 29 years without developing a leak in the city block containing the specimens. This block also contains five services, one of which served without a leak for more than 25 years. Another service developed a leak in 9 years. Using Scott's pit-depth—area relation, the average of the maximum pit depths for 500 feet of 6-inch cast iron is 468 mils for a period of 29 years. This indicates that the cast-iron pipe in this soil, the wall thickness of which is 430 mils, should have developed, on the average, one puncture per block at the time the information was obtained. However, the comparison of the field and experimental data is of doubtful value because the extent of

TABLE 52.—Depth of the deepest pit on 4 feet of unevaled line pipe and on the adjacent 3-inch control pipe.

[Depth in mils]

								Ž.	Zilena						
		I.	Ħ	H.	IV.	ν.	VI.	VII.	VIII.	X	×	XI.	XIII.	XIV.	XVI.
	Specimen	Temple, Tex.	Arkansas City, Kans.	Beau- mont, Tex.	League City, Tex,	Preble, Ind.	Council Hill, Okla,	Сапеу, Капя,	Top (fully, Tex.	Long Beach, Calif.	Mt. Auburn, Ill.	Skintook, Okla.	Bunkie, La.	Cham- bers- barg, Pa.	Cholame Flats, Calif.
	Control	36 40	47	235 IIb	20 52	88	50.02	42	118	110	88	130	\$8	\$3	
<u> </u>	(Line	24 36	57 39	280 H	73	55 29	02 02 03	69	+ 808	100	24.	159	23.5	<b>4</b> 5	
·	[Line	30 43	88	216 H	60 48	65 34	42 65	46 51	105	78 135	45 49	105	328	45	
	Control	36	89 :	229	51 70	41	30	47 50	158	47 51	62 75	109	126 32	<b>%</b> \$	
, U	Control	25 61	<b>4%</b>	160	75 68	45	328	41 56	135	72	88	98		31	:
10	(Line	79	65	14t H	40 60	58 40	43	94	199+	100 105	99	75	150 27	: 23	
~0	(Line	36 49	65 40	200	68 50	45 40	41 37	55 51	168	83 109	57	120+ 49	122 38	26 43	
-0	(Line.	26 48	65	216	283	35 42	46	99	190	809	46 57	92+	109	37	
-0	(Line, Control	35 60	37	183	44	36	57	116 95	87	83	56 68	102+	424	38	
-0	(Line	30	58	126	70 120	41 37	53	177 45	121+	154 89	44 51	+ 201	62	46	
-0	(Line	54 49	68 46	200	88	45	51 52	61 50	156+	135 89	48	51+	76	41	
-0	Control	58	45	131 152	202	46 59	53	76 63	157+	175 98		102+	105		
101	Control	38	61 65	127 123		50 38		106		60 51	75	52+	152		

the pipe in Ramona loam has not been determined, also as the pitting factor for cast iron is greater than for steel, the value for a should be less, and furthermore, cast iron may be punctured by corrosion and still

carry low-pressure gas without leaking.

A little additional light on the relation between the corrosion of pipe lines and that of short lengths of pipe is furnished by the American Petroleum Institute's tests of protective coatings [61]. Bare sections of pipe line were left between two coated sections. Opposite each bare section a 30-inch length of 3-inch pipe was placed in each trench as a control specimen. Table 52 shows the deepest pit on sections of the line and adjacent control pipes. The line pipes differed in diameter.

Although the area of the pipe was from five to nine times that of the control, there is no definite relation between the pit depths on the two areas. Because some of the coatings on the line were rather porous, the pipe beneath them may have been anodic on account of differential aeration, and so may have protected the bare sections. The 13 sections were distributed along about 1,100 feet of pipe, and the difference in pit depths on different sections illustrates the dispersion of pit depths on pipe lines. However, table 28 is a much better presentation of this

phenomenon, as it pertains to a pipe that was not coated.

Although a review of these comparisons of test data with actual performance indicates no very close relation between the two, usually the soils which the test has shown to be corrosive are shown by experience also to be corrosive. Corrosion is affected by so many factors that it can scarcely be expected that any test could completely duplicate the conditions which buried metals encounter in service. Knowledge of service conditions is in most cases so incomplete that it would be difficult to use an equation that would take account of all significant field conditions even if such an equation could be developed.

# VII. RESULTS OF FIELD TESTS ON MISCELLANEOUS NONFERROUS MATERIALS

## 1. COPPER AND COPPER ALLOYS

The first copper-alloy specimens buried in 1924 in the National Bureau of Standards tests consisted of four high-copper cast brasses connected as caps to nipples of brass (Muntz metal), lead, and galvanized steel. The final results of these tests are reported in Research Paper RP945 [62]. The cast caps each weighed about 150 grams (5.3 ounces). The nipples were cut from ½-inch pipe, each having an exposed length of 2-inches, that is, about 5 square inches. Table 53 shows the approximate composition of the cast caps.

Mixture No.	Cu	Sn	Pb	$\mathbf{z}_{\mathtt{n}}$	Fe
1 2 3 4	85.5 87.0 91.8 75.0	3.5 5.9 0.1 1.4	4.8 1.8 0.3 2.6	6.2 5.3 7.7 20.0	1.0

Table 53.—Composition of cast brass caps.

The small size of the specimens, their irregular shapes, and the galvanic potentials and small corrosion losses combined to make precise results impossible and a detailed report on the specimens from each site seems unnecessary. Figure 22 shows the average rates of loss for

⁵ See section VIII, B for a description of the tests.

each material on all soils. Figure 23 permits a comparison of these losses with losses of similar materials unaffected by galvanic action. It was evident that the connecting of the different materials accelerated the corrosion of the nipples, and reduced the corrosion of the cast caps. The intensity of the galvanic corrosion was affected by the relative areas of

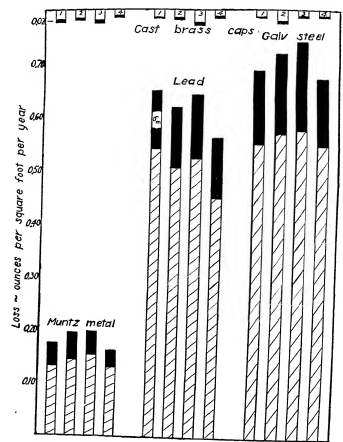


FIGURE 22.—Average rates of corrosion of cast-brass fittings and attached nipples in 15 soils.

the anodic and cathodic parts of the combination and no generalization could be drawn, except that connecting two metals in the presence of an electrolyte tends to cause one to corrode more rapidly and reduces the corrosion of the other.

In 1926 a few copper and copper-alloy pipes and rods were buried in 47 soils. The compositions of these materials are given in table 9. Table 54 shows the rates of corrosion of these materials based on exposures of approximately 8 and 13 years. As stated in the discussion of the ferrous

materials, rates of corrosion change with the duration of the exposure and should not be used to predict corrosion for any period except that used in deriving the rate. However, the data in table 54 are comparable with each other. They show that in most soils the losses of weight during 8 years were very small, and that the maximum penetration was slight.

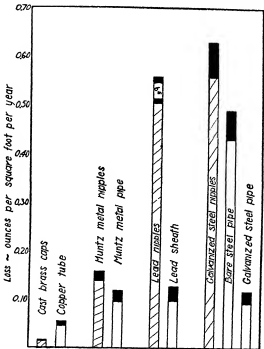


FIGURE 23.—Effect of interconnecting different metals.

The brass caps were connected to lead and galvanized iron. All the nipples were connected to brass caps.

The table puts the brass alloys in too favorable a light because it does not show the deterioration of these materials due to dezincification or selective corrosion. This is recognized as copper-colored spots or areas on the surface of the metal and by a loss of strength, but not of form, if the dezincification penetrates deeply. According to McKay and Worthington [63], accidental variations in the soil solution or in the alloy result in concentration or galvanic cells and at the anodic area the alloy goes into solution, copper is then redeposited from the solution at or near the points where the corrosion occurred. This deposition of copper sets up another galvanic cell and corrosion continues to spread and to penetrate the alloy. The acid around the anode keeps the zinc in solution and it migrates into the soil. According to these authors, the most favorable conditions for dezincification are a good conducting solution and a slightly acid condition, with the presence of oxygen. In the Bureau's tests the most serious dezincification occurred in einders that had a pH of 8.0 and poor aeration. Serious dezincification also occurred in Docas clay, a poorly-drained soil with a pH value of 8.3. Table 54 indicates that some dezincification of three high-zinc brasses occurred in all soils in the tests of corrosion-resistant materials. Figures 24 and 25 are photomicrographs of corroded areas of two types of brass showing the uncorroded metal and the dezincified (corroded) metal.



FIGURE 24.—Yellow brass exposed 9.53 years to Sharkey clay.

Transverse section showing uncorroded metal (below) and dezincified metal (above) × 500.

Tables 55 and 56 show the losses of weight and pit depths for a variety of copper alloys exposed for approximately 9 years to 13 soils that were corrosive with respect to ferrous materials. The corrosion of low-carbon steel in these soils is shown for comparison. The relative corrodibilities of the several alloys with respect to each other remain approximately the same except in a few soils.

The corrosion resistance of copper has been attributed [20] to its nobility, that is, low solution potential and to the density of the oxide film which forms on the surface. Sulfide and reducing conditions are corrosive to copper. It will be noted that the soils that are most corrosive to copper are mostly wet, organic soils. Such soils frequently contain bacteria which reduce sulfates to sulfides. According to Ewing [29], the corrosion of copper in such soils as Docas clay is probably caused by differential aeration, and is stimulated by the action of chlorides. The same author attributes the corrosion of copper in such soils as muck and peat to organic acids that tend to dissolve the protective oxide film. The most corrosive condition in the Bureau's test sites was in cinders (soil 67). As an analysis of a sample of these cinders from which the large clinkers had been removed showed 26 percent of carbon, it seems probable that much of the corrosion was the result of galvanic action between the carbon and the test specimens. Sometimes, cinders from completely burned coal are not corrosive. They may be inert and act much like sand. To permit a better comparison of the materials, table 57 has been prepared. It is based on the averages of data for four periods of exposure in 11 of the soils of table 56. Data for soils 63 and 67 were omitted because of the destruction of some of the materials at these test sites.





FIGURE 25.—Two-and-one leaded brass (leaded silicon brass) exposed 9.53 years to Sharkey Clay

A, transverse section showing transition from uncorroded metal at left to complete dezincification at right,  $\times$  15: B, section showing partial dezincification, the dezincification being more severe at the left,  $\times$  250.

TABLE 54.—Corrosion of copper-rich alloys exposed for 8 years and for 13 years.

[For the condition, the letters indicate the worse of 2 specimens. The figures are rates of maximum penetration, in mils per year.]

P. Definite pritting, no pils greater than 6 mils.

U. Unaffected by corresion, impossible to measure penetration because of even destruction of surface.

D. selective corresion, impossible to measure penetration because of even destruction of surface.

D. selective corresion such as desiredification over large areas.

E. destruction such as poise.

E. destruction is spoise.

E. destruction to provide a proper and the destruction of surface.

E. destruction by destruction for the provident and the property of the provident and the providen

			Ачегадо	rate of loss	Average rate of loss of weight (02/ft?)/yr	ft²)/yr		· de la	Condition o	Condition of surface and pit depth	pit depth	
Soil No.	Duration	Copper pipe	r pipe	Br	Brass	Cu-Zn-Ni	Cu-Al	Copper phys	pipe	Bruss	Cu-Zn-Ni	Cu-AI
	of test	M	Ъ	Pipe, B	EII. Me	< <	z	M	A	B B	Just,	, z
H6360 410	Year's 8.07 13.54 8.01 7.94 13.36	0.060 .023 .027 .019	0.003 .016 .029 .019	0.097 .065 .074 .144	0.185 .090 .109	0.157 .101 .156 .176	0.057 .019 .063 .070	1.5 1.3 0.6	HEARA	M. 1.7.7. 1.7.7.4. 1.0.0.4.	5,4,1,1,0 0,9,8,7,0,0 0,6,8,7,0,0 0,0,0,0,0,0 0,0,0,0,0,0 0,0,0,0,0	MACG4
8.5°8°4°8°4°8°4°8°4°8°4°8°4°8°4°8°4°8°4°8°	13.33 13.44 17.96 13.44 13.18	.011 .036 .024 .030	.0093 .026 .019 .036	.017 .188 .018 .174 .235	.319	.056 .155 .136 .165	.0023 .032 .026 .011	P P.5 0.8	P. 1.3	P, d M, d 0, 5, D 0, 6, D	P, D 0.07 1.5, d 0.9, D 0.8, D	Р Р Р Р
12 13 14 15 16	7.96 7.96 7.97 7.97	.312 .023 .040 .013	.278 .031 .025 .016	222 101 108 060 170 170	.024	.199 .012 .096 .045	.090 .028 .017 .026	1.5 P P 5 3.3	ж М 1.9	P, D M, D P, d 1.3, D	M, d M, d 1.9, d d.9, d	a, M. A, a a
17. 18. 19. 20. 22.	7.96 7.97 7.97 8.08 7.98	.037 .0076 .039 .042	.040 .0077 .040 .039	.059 .021 .122 .044 .195	.257 .070 .067 .198	.108 .044 .165 .138	.049 .013 .027 .044	M U 1.1 1.5	MM P P P P P P P P P P P P P P P P P P	M, d M, d P, D P, d 1.1, D	9.4.1.4.4 2.8.2.2.2.2.2.2.2.2.2.2.2.2.2.2.2.2.2.2	MOMA M, d M, D
23 24 25 27	7.96 13.18 7.96 13.44 13.58	.118 .019 .012 .013	.135 .018 .011 .012	.752 .025 .051 .072 .046	1.85 0.142 .058 .080	.288 .030 .126 .018	.160 .043 .034 .024	1.9 PP PP PP 9	E DAG	Z P,P,d P,D	PFI-PG PDGD	APDPB

d M, d d MP, d	M, d P M, d	ւ թ. 2.3	PD'd
P, d 1 9, d P, D P, D 1.3, d	1.3, d P, d 1.0, d 9.7, D	P, d 1.9, d 0.5, D U, d	2.4, d 1.0, d P, D
M, D P, D P, d	R, D B, D D, 7, D P, D	M, d 1.8, D P, D M, d	1.3, D d P, D
PUPPE	1.0 P P 1.3	P.1.1	P 1.0 0.8
AFFFD	0.9 M P P 1.8	M2.742	P 1.0 1.2
.107 .082 .0071 .026	.040 .021 .039 .014	.019 .092 .035 .057	.026
. 163 . 257 . 032 . 022	.208 .177 .063 .087	.034 .385 .126 .154	.248 .081 .101
	.218 .080 .422 .039	.037	.214
.080 .275 .014 .019	.175 .057 .020 .049	.028 .349 .083 .095	175 016 .037
.079 .116 .0097 .0086	.117 .022 .016 .019	.043 .168 .030 .049	.080
.084 .123 .0078 .0083	.137 .016 .017 .019	.025 .125 .027 .047	.079 .033 .032
7.96 7.97 13.39 13.66 7.93	7.96 7.97 7.95 13.61 7.97	7.97 7.98 13.38 7.97 7.95	7.97 7.95 13.36
28 29 30 31 32	33. 35. 37.	38. 40. 42. 43.	44. 45. 47.

TABLE 55.—Loss of weight of copper and copper alloys exposed for 9 years.

In ounces per square foot]

	Alloy	Sa Muntz 97% Cu 88% Cu	I, E N "N, D G N	0.41 0.52 0.37 0.30 0.36 0.27 4.09 3.33 0.00 0.27 4.09 0.00 0.00 0.27 0.28 0.20 0.33 0.00 0.00 0.00 0.00 0.00 0.00
	_	l- Brass ed 66% Cu s 33% Zn	f	
		- 2-and- 1-leaded l brass	M	0 0 47 0 47 0 47 0 90 0 100 0 10
		Admi- ralty metal	Ħ	0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000
-		Red	F	0.25 0.25 0.25 0.12 0.12 0.13 1.75 0.14 0.25 0.14 0.12 0.13 0.14 0.13 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14 0.14
		Deoxi- dized copper	Ą	0.22 .18 .2.78 .2.78 .0.11 .4.22 .3.4.22 .3.4.22 .3.4.22 .3.6.24 .1.50
		Fough pitch copper	۵	0.24 1.97 1.97 1.97 1.93 1.38 1.38 1.38 1.38 1.38 1.38 1.38 1.3
		Ex- posure		Years 9.47 9.41 9.11 9.51 9.24 9.53 9.25 9.25 9.25 9.25 9.25 9.26 9.27 9.28
	Soil	Tvne		Cecil clay loam   Pears   0.24   0.25   0.25   0.25
		No.		255 255 255 255 255 255 255 255 255 255

Table 56.—Maximum penetration of copper and copper alloys exposed for 9 years.

In mils

M, shallow metal attack, roughening of the surface but no definite pitting. P, definite pitting, no pits greater than 6 mils.

s, uniform correspon, impossible to measure true penetration.
D, selective corresion, such as destinction over large meas.

d, selective corrosion over small areas. Z, specimens destroyed by corrosion (dezincification). +, one or both specimens punctured.

	Low-carbon steel	×	59 59 110, 8 27, 8 27, 8 696 87 154+(5, 7) 1134+(2, 7, 7) 134+(2, 5, 7)
	75% Cu 20% Ni 5% Zn	G	8,0,0,0,0,0 0,0,0,0,0,0,0,0,0,0,0,0,0,0,
Alloy	95% Cu 3% Si 1% Mn	Q	888 M M M M M M M M M M M M M M M M M M
N. A.	98% Cu 1.5% Si 0.2% Mn	N q	8 49 PP s 0,08
	98% Cu 1.5% Si 0.2% Mn	Z	PP
Bronze	97% Cu 1% Si 1.8% Sn	Ħ	202 122 202 202 202 237 222 222 102 102
	Muntz metal	L	6, D 6, D 6, D 6, D 7, D 7, D 7, D 8, D 8, D 8, D 8, D 7, D 8, D 8, D 8, D 8, D 8, D 8, D 8, D 8
F	66% Cu 33% Zn	J	6, D 16, D 8, D 8, D M, D M, D 13, D 12, D 12, D 12, D 12, D 12, D 13, D
	and-one leaded brass	М	7.7.7.00 6.7.7.00 6.7.00 6.7.00 7.7.00 7.7.00 7.7.00 7.7.00 7.7.00
er targe areas	Admiralty brass	н	20, D 20, D 20, D 4 P, D 4 P, D 7 P, L 7 P, D 68, S, D 68, S, D 68, S, D
neartion ov	Red	뚄	10,77,00 10,00 10,00 14,70 14,00 16,40 17,00 16,40 16,40 16,40
Selective corrosson, ston as dezinchoation over large areas,	with soldered joints	8 M	113 4 15 118 118 110 110 118 118 118 118
Troston, st	dized	A	9.14 9.14 9.8 10,8 10,8 10,8
Tours.	pitch copper	Ö	6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6
P.	Soil No.		55 55 55 55 55 55 55 55 55 55 55 55 55

* These specimens had streamlined caps and couplings soldered in place.

* These specimens had haved joints; dath for only I specimen.

* Maximum pits for individual specimens differed from each other by more than

50 percent.

4 Data for 3 specimens.

A number in parentheses after the pit depth indicates that 1 or both specimens of a number in parentheses after the pit depth indicates that the specimen was punctured a previous removal was punctured, e. g., (5) indicates that the specimen was punctured after 5 years.

s Data for I specimen only.

Halo in I specimen due to dezineification. The mate, although it had no measurable pits greater than 8 mils, reads bely dezineified, as indicated by the flat sound when the pipe was struck with another piece of negal.

I specimen destroyed by dezineification. Average pit depths of 1939 removals are greater.

	Material		Compositio	on		Stand-	Stand-
Symbol	Туре	Cu	Zn	Pb	Aver- age	ard devia- tion	ard error
A Dec F Red H Adr K Two J Bra	igh-pitch copperxidized copperbt brass hiralty metal brass ss.	99 9 99 94 85 2 71 3 67.1 66.5 60.1	14 8 27.4 31.1 33.1 39 6	0.8 .4 .4	700 144 105 121 176 225 492	% 93 32 53 176 229 942	% 
N Cop	nze per-silicon alloydo.	97 2 98.1 95.5	Si 1.0 1.0 3.2 Zn	Sn     1.8	193 154 142	144 79 63	21 11 9
G Cop	per-nickel alloy	74.5	5.0	20 0	104	54	8

Table 57.—Relative loss of weight of copper and copper alloys.

[Average of four periods of exposure, 11 soils]

The high standard errors for some materials indicate that their performance relative to copper was different at different test sites or for different periods of exposure. The table should be used with care, as the performance of a material under some specific conditions may be quite different from the average of its performance under a number of conditions. For example, Muntz metal corroded very little in tidal marshes, soils 43 and 63, but was quite unsuitable for most other corrosive soils.

The copper alloy containing approximately 60 percent of copper and 40 percent of zinc, commonly known as Muntz metal, is subject to dezincification under several soil conditions. To determine whether or not the addition of arsenic to this alloy would prevent dezincification, specimens of Muntz metal plates containing 0.08 percent of arsenic were added to the tests in 1939. Unfortunately, similar plates of ordinary Muntz metal were not buried at the same time. This makes it necessary to compare the behavior of the new materials with that of a section of pipe buried 7 years previously. Obviously, under these conditions small difference in performance may be accidental. Table 58 shows the loss of weight and the condition of the two materials exposed for approximately the same periods. It appears that the addition of 0.08 percent of arsenic was insufficient to prevent dezincification of the brass.

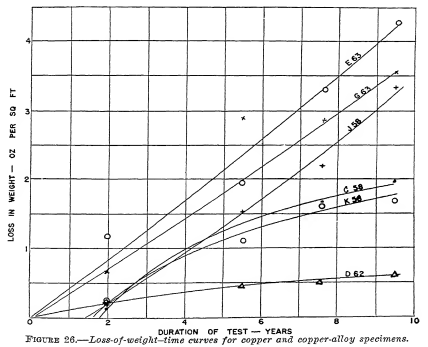
The effect of time on the rate of corrosion of nonferrous metals has not been examined as thoroughly as was done for ferrous materials. The loss of weight and pit depths in most soils are small, and this makes their accurate determination more difficult. In addition, the copper and copper-alloy specimens in the Bureau tests were exposed for shorter periods than the ferrous materials and the number of examinations has been less. In figure 26 are shown the losses of weight per unit area for several materials in three soils. The letter identifies the material (see table 57) and the number relates to the soil (table 55). The curves were drawn arbitrarily and might have been drawn differently, possibly with different significance. As drawn, curves J58, C58, and K58 indicate that the materials did not begin to corrode for a considerable time after burial, possibly because of a temporarily protective oxide film.

Table 58 .- Loss of weight and maximum penetration of Muntz metal exposed 2 years.

M. shallow metal attack, roughening of the surface but no definite pitting.
P. definite pitting but no pits greater than 6 mils.
s. uniform corrosion, no reference surface.
D. selective corrosion by many the cities over large areas (several square inches per square foot).
d. selective corrosion over small areas.
Z. specimens destroyed by dezincification.

	Soil	Ar	senical M (sheet	untz metal ) Bı	Mu	ıntz metal	(pipe) L
No.	Type	Expo- sure	Loss of weight	Maximum penetration	Expo- sure	Loss of weight	Maximum penetration
53 55 56 58 60	Cecil clay loam. Hagerstown loam. Lake Charles clay. Muck. Rifle peat.	2.03	oz/ft ² 0.18 .16 .55 .54 1.87	Mils P,D P,d 12,D P,D 13,D	Years 1.96 1.89 1.99 1.99 1.99	oz/ft ² 0.19 .19 .14 .20 1 73	Mils P,d 6,d M,d P,d 9,d
61 62 63 64 65	Sharkey clay. Susquehanna clay. Tidal marsh. Docas clay. Chino silt loam.	1 90 1.88	0.40 .32 a 036 .47 .21	P,D P,D *P P P,D	0.95 1.93 2.04 1.91 1.91	0.16 .33 1.41 4 02 2.43	P,d P,d M P,D M,D
66 67 69 70	Mohave fine gravelly loam Cinders Houghton muck. Merced silt loam	1 86 1.90 1 90 1.90	.30 15.25 0.30 .28	8,d 122,s,D P,D 12,D	1.92 2.02		^b P,d Z

Data for 3 specimens.
 Data for 1 specimen.



See tables 55 and 57 for identification of soils and specimens.

As in the case of the ferrous materials, the pit-depth data for the copper alloys are quite erratic and, because of the smaller depths and fewer removals, it is quite impossible to plot satisfactory pit-depth-time curves from most of the data. Some of the most nearly consistent data are shown in figure 27. The curves indicate that there is only a slight

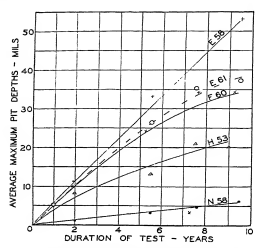


Figure 27.—Pit-depth-time curves for several copper-alloy specimens.

See tables 55 and 57 for identification of soils and specimens.

tendency if any for the rate of corrosion to decrease with time. This is probably because of the solubility of the corrosion products.

Somewhat more definite conclusions can be drawn after the specimens now in the ground have been removed, but it is improbable that the exact rates of corrosion of the copper-alloys can be determined from the tests. It is evident that, with a few exceptions with respect to soil types and alloys such as Muntz metal, the deterioration of copper alloys exposed to soil is much slower than that of any of the ferrous alloys except those high in silicon, chromium, or nickel. If more specific information is desidered, new tests in which soil and other conditions are controlled much more closely and which include a much greater number of specimens of the same material in each test site must be undertaken.

#### 2. LEAD

Lead is used extensively underground in the form of water service pipes and cable sheaths. Most cable sheaths are placed in fiber, cement, or vitrified-clay ducts and are in contact with soil only as it is washed in from manholes. Occasionally, lead cable sheaths are laid directly in the ground. Parkway and similar power-distribution cable is laid directly in the earth, but in such cable the lead is surrounded by wrappings of steel tape and jute or other fiber impregnated with a bituminous material.

Table 59 shows the loss of weight and maximum depth of penetration of antimonial lead and commercial lead cable sheaths exposed for from 10 to 17 years. For comparison, similar data are given for a 12-inch

length of 3-inch Bessemer steel pipe (113 square inches) in the same soils. The area of the antimonial lead sheath was in most cases 370 square inches, whereas that of the commercial lead was 150 square inches. The duration of exposure of the lead and steel specimens in table 59 was somewhat different in some soils.

Table 59.—Loss of weight and maximum penetration of lead cable sheath exposed 10 to 17 years.

	Soil	Dura-		nial lead A		rcial lead H		ner steel M
No.	Туре	of expo- sure	Loss of weight	Maxi- mum pene- tration	Loss of weight	Maxi- mum pene- tration	Loss of weight	Maxi- mum pene- tration
1 2 3 4 5	Allis silt loam. Bell clay. Cecil clay loam. Chester loam. Dublin clay adobe.	Years 11.65 a15.48 a10.02 12.00 a15.56	oz/ft ² 4.02 1.56 0.63 1.79 3.51	Mils 120+ 45 8 64 120+	oz/ft ² 2.01 1.20 0.59 2.13 6.66	Mils 112+ 30 18 51 112+	oz/ft ² 11.56 7.12 4.34 6.92 8.58	Mils 101 58 72 84 62
6 7 8 10 11	Everett gravelly sandy loam Maddox silt loam Fargo clay loam Gloucester sandy loam Hagerstown loam	*15.53 16.94 11.76 11.95 11.92	0.37 1.47 0.73 .89 .48	22 39 12 26 36	0.26 1.60 1 0.38	28 32  15 15	1.88 5.06 8.29 4.94 1.74	20 48 111 54 81
12	Hanford fine sandy loam	*15.59	1.88	30	1.85	43	5 32	86
14	Hempsted silt loam	11.76	2.13	68	0.57	24	4 58	95
15	Houston black clay	*10.06	0.52	16	.36	25	8.11	72
17	Keyport loam	11.78	.43	36	.33	20	9.65	48
18	Knox silt loam	11.71	.50	20	.18	10	2.40	44
19	Lindley silt loam	11.63	1.31	36	.46	15	3.32	60
20	Mahoning silt loam	11.65	3.54	78	3.12	51	5.99	56
22	Memphis silt loam	11.65	1.72	32	1 00	19	7.78	65
24	Merrimac gravelly sandy loam	11.95	0 25	18	0.18	19	1 40	30
25	Miami clay loam	11.67	.53	44	.33	28	3.31	57
26	Miami silt loam	11 52	.47	22	.21	29	3.77	38
27		*15.69	1.37	39	.67	31	8.86	84
28		9 60	2.12	52	.66	10	15.54	152
29		*10.08	3.55	14	3 45	34	14.84	119
30		17.04	2.28	56	1.04	51	6.91	76
31	Ontario loam	*15.73	0.37	<6	0.28	15	4 08	66
32		11.66	.59	12	.33	18	4.15	62
34		12.00	3.12	120+	.81	112+	5.31	50
35		*15.59	0.19	12	.31	37	3.21	36
36		*15.69	.69	17	.48	22	2.58	49
38	Sassafras gravelly sandy loam.	12.01	.38	14	.30	25	2.23	43
39	Sassafras silt loam.	12.00	1.77	47	1.18	36	6.36	94
41	Summit silt loam.	17 41	0.77	41	0.50	27	7 03	101
43	Tidal marsh	12.02	.51	28	.23	13	13.15	75
44	Wabash silt loam.	11.61	.82	22	.44	13	3.39	83
45	Unidentified alkali soil	11.73	.45	19	.24	23	9.29	98
46	Unidentified sandy loam	12.00	.98	42	.18	14	4.81	114
47	Unidentified silt loam	17.43	1.12	32	.79	30	5.37	<b>4</b> 8

a Duration of exposure for the Bessemer steel pipes was approximately 2 years longer.

Figures 28 and 29 show the progress of the loss in weight for a few soils. The letters near the curves indicate the kind of lead tested, and the numbers are the numbers of the test sites. The two identifications permit the reader to determine from tables 10 and 20 the known conditions of the test. Four of the curves indicate that the rate of corrosion of the lead specimens increased as the specimens became older; two other curves show decreasing rates of corrosion. All of the soils involved are nearly neutral in reaction. The curves do not indicate the distribution of the corrosion. Some idea of this can be gained from table 59. The pitting was greatest in soil No. 1 and least in soil No. 8, which contained some sulfates. After examining the Bureau's data on com-

mercial lead specimens exposed from 10 to 17 years, Ewing [29] divided the types of corrosion as follows: No corrosion, concentrated attack, and uniform attack. The soil characteristics he associated with the first class were high pH and high sulfates. The second type of corrosion occurred

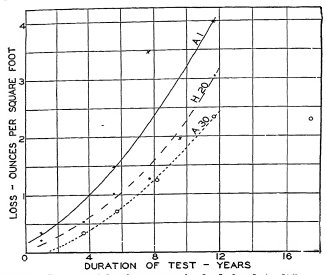


FIGURE 28.—Loss-of-weight-time curves for lead sheath in different soils.

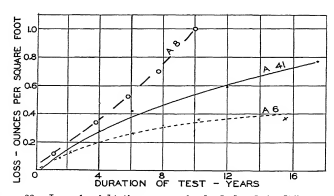


FIGURE 29.—Loss-of-weight-time curves for lead sheath in different soils.

in neutral soils, where apparently the original protective film broke down in spots. The third class of corrosion occurred in reducing soils, which apparently destroyed the protective film. Ewing recognized that other factors affected the corrosion of many specimens, but the data are insufficient to identify these factors.

In 1937 three varieties of lead-alloy pipe, such as are used for water service, were buried in 14 soils. Table 60 shows the extent of the cor-

rosion of these materials after exposures of approximately 4 years. No material is either worst or best in all soils. On the average, the antimonial lead is least subject to pitting. However, in the earlier tests. antimonial lead that contained much less antimony than the later antimonial-lead specimens was inferior to commercial lead. The differences in losses or pit depths are small, and may be due to chance. More specimens remain to be removed at later dates. A point shown quite positively is that in many soils the pitting of lead is sufficiently deep within a few years to puncture lead coatings of the usual thickness. The potential between lead and iron accelerates the corrosion of iron when the two metals are in contact in the presence of moisture. The data indicate, therefore, that in many soils only a very thick coating of lead would protect iron, and even such a coating would be unsatisfactory in a few soils where the rate of penetration for lead is as great as it is for iron.

Table 60 .- Loss of weight and maximum penetration of lead pipe exposed 4 years.

	Soil	Chemic	al leada	Telluriu	m leadb	Antimon	ial leade
No.	$_{\rm Type}$	Loss of weight	Maxi- mum pene- tration	Loss of weight	Maxi- mum pene- tration	Loss of weight	Maxi- mum pene- tration
53 55 56 58 60	Cecil clay loam Hagerstown loam Lake Charles clay Muck Rifle peat	$0z/ft^2 \ 0.21 \ .20 \ .45 \ 2.41 \ 0.28$	Mils 12 26 37 28 15	$0z/ft^2$ 0.31 .28 .82 2.80 0.20	Mils 20 26 48 56 10	$0.2/ft^2 \ 0.22 \ .15 \ .50 \ 2.12 \ 0.22$	Mils 10 18 52 58 dP
61 62 63 64 65	Sharkey clay	2.21 0.93 .015 .19 .13	39 29 18 16 24	1.75 0.64 •.015 .18 .16	30 31 e12 11 16	1.75 1.03 0.013 .19 .21	42 30 16 12 15
66 67 69 70	Mohave fine gravelly loam Cinders Houghton muck Merced silt loam	$^{.10}_{12.21}$ $^{0.81}_{.12}$	34 104 15 14	.12 13.22 1.08 0.15	41 94 12 27	$\begin{array}{c} .12 \\ 4.21 \\ 1.04 \\ 0.14 \end{array}$	15 90 7 12

#### 3. ZINC AND ALUMINUM

Zinc is not used extensively for underground pipes or containers but is frequently used as a coating and in recent years it has been used in considerable quantities as anodes for cathodic protection. For the latter use it is essential that it continue to corrode as it is the current associated with the corrosion of the zinc that protects the metal to which zinc is connected.

Table 61 indicates the extent of the corrosion of two varieties of zinc plates exposed for approximately 4 years. Comparison of this table with the data on 2-year exposures [64] shows that in most soils the corrosion was nearly proportional to the duration of the exposure. However, in four soils high in salts the rate of corrosion was much less for the longer exposures. Comparison of the rates of corrosion of zinc and steel shows that steel loses weight from two to four times as fast as zinc in most soils during the first 4 years of exposure. The maximum penetration of

a Cu, 0.056%; Bi, 0.002%; Sb, 0.0011%. b Cu, 0.082%; Te, 0.043%; Sb, 0.0011%. c Cu, 0.036%; Bi, 0.016%; Sb, 5.31%. d P, definite pitting but no pits greater than 6 mils. e Data for 1 specimen only.

steel is correspondingly greater for 4-year-old specimens. However, as the rate of corrosion decreases more rapidly for steel than for zinc, the ultimate rates for the two materials may be nearly the same for long periods of exposure. In some of the highly organic soils, zinc loses weight

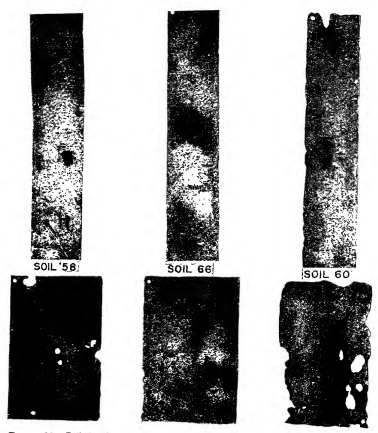


FIGURE 30.—Rolled (above) and die-cast (below) zinc buried in muck (soil 58), in Mohave fine gravelly loam (soil 66), and in Rifle peat (soil 60), approximately 4 years each.

faster than steel, and the penetrations of the two metals are of the same order. Figure 30 shows specimens of rolled and die-cast zinc exposed to three soils.

Figure 31 shows the relation of the average maximum penetration to the duration of exposure of zinc specimens to five soils. Each point is based on the average of the deepest pits on each of two specimens. In two of these soils the rates of penetration appear to increase with the period of exposure.

Table 61 .- Loss of weight and maximum penetration of zinc plates exposed for 4 years.

	Soil	Rolled	zinc, Z	Die-cast	zinc, CZ
No.	Туре	Loss of weight	Maximum penetration	Loss of weight	Maximum penetration
53 55 56 58 60 61 62 63 64 65	Cecil clay loam Hagerstown loam Lake Charles clay Muck Rifle peat Sharkey clay Susquehanna clay Tidal marsh Dogas clay Chino sitt loam	3.42 5.09 10.36 0.96 1.24 b2.30	Mils 10 a 8 b26 66 a100  8 9 34 18 36	oz/ft² 0.54 .61 4 96 6.33 14.98 1.12 0.60 1.43 2.53 0.76	Mils 22 20 30 c125+(2) 125+ 28 16 24 20 16
66 67 69 70	Mohave fine gravelly loam. Cinders. Houghton muck. Merced silt loam.	^b 2.61 d12.16 1 70 d1.62	^{b28} ^{a118} +(2) ¹⁰ ^{b102} +	4.74 13.08 1.64 42.19	124 + 125 + 36 580 +

The National Bureau of Standards obtained a few data on aluminum and two of its alloys exposed to only five soils. The test specimens measured 2 by 6 inches. In some of the soils, the duralumin was completely converted to a greenish-white paste. Intergranular corrosion raised ridges and blisters, beneath which was a white powder on some of the specimens. The unalloyed specimens were the best of the group. Table 62 shows the loss of weight and maximum penetration of the thin aluminum specimens, exposed approximately 10 years, and similar data on zinc and iron for comparison. None of the thin materials was satisfactory for use unprotected in the corrosive soils to which they were exposed.

Table 62.—Loss of weight and maximum penetration of zinc and aluminum exposed for 10 nears.

				,0, 10	goard	•					
Duration of exposure (years)		10	. 16	10	.08	10	.05	10	.73	10	. 55
		Soi	l 13	Soi	1 29	Soi	l 42	Soi	1 43	Soil	l 45
25	Iden-	fines	rd very andy am	Mı	uck	Susque	ehanna lay	Tidal	marsh		ntified fali oil
Material	tifica- tion	Loss of weight	Maxi- mum pene- tra- tion	Loss of weight	Maxi- mum pene- tra- tion	Loss of weight	Maxi mum pene- tra- tion	Loss of weight	Maxi- mum pene- tra- tion	Loss of weight	Maxi- mum pene- tra- tion
Sheet zinc. Cast zinc Sheet zinc Aluminum Al-Mn-alloy Duralumin Open-hearth iron Steel +0.2% Cu	Z1 Z2 P C1 C2 C3 A	oz/ft² 3.52 3.47 1.89 0.086 .38 D 9.92 D	Mils 40 71 53 21 45+ D 125+ 62+	oz/ft ³ 4.66 5.37 3.85 D °0.97 D 5.86 6.91	Mils 39 55 62+ 62+ 62+ D 62 62+	oz/ft ² 0.85 .79 1.06 0.35 .20 1.39 5.61 5.40	Mils 17 18 20 62+ 14 62+ 70 59	oz/ft² 2.48 2.38 4.85 0.18 2.22 .15 D	Mils 36 73 40 <6 13 <6 125+ 62+	oz/ft² aD 9.93 D 0.49 .33 c.56 D D	Mils 62+ 104 62+ 46+ 20 62+ 125+ 62+

D =destroyed by corrosion.

a Uniform corrosion; no reference surface left.
b Data for individual specimens differed from the average by more than 50 percent.
- Hndicates that 1 or both specimens punctured by corrosion from 1 side of the plate. (2) indicates that 1 specimen from the previous removal was punctured after 2 years.
d Data for 1 specimen only; the other specimen was destroyed by corrosion.

b +=1 or both specimens punctured because of corrosion.
 Data on 1 specimen only. The other specimen was destroyed by corrosion.

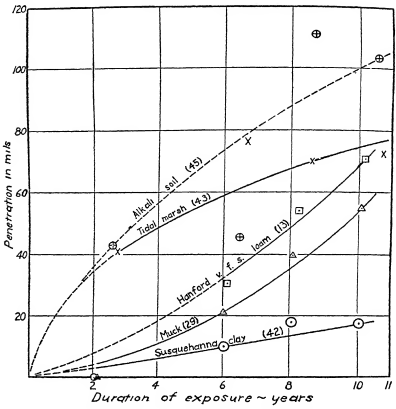


FIGURE 31.—Relation of the maximum penetration of zinc to the duration of exposure.

## 4. ASBESTOS-CEMENT PIPE

The use of asbestos-cement pipe began in Italy about 1916. Since that time it has been used more or less extensively in many constructions in Europe and to a considerable extent in Japan. So few data on the effect of soils on this material are available that the Bureau's 1941 report is reproduced here in full.

Asbestos-cement pipe is a mixture of asbestos fiber and cement. The pipe is built up by a continuous process on a revolving steel mandrel. This is a followed by a curing process. Because asbestos-cement pipe is nonmetallic, it is, of course, not subject to galvanic corrosion, tuberculation, or electrolysis. Pipe of this type is manufactured largely for use in transmission mains and services where the working pressure ranges from 50 to 200 pounds per square inch.

To determine whether or not asbestos-cement pipe is subject to deterioration under soil-exposure conditions, specimens were buried at 15 test sites in 1937. The specimens were 12 inches long and 6 inches in

diameter and had an average wall thickness of 0.72 inch. These specimens were cut from class 150 pipe. Two specimens were removed from each test site at each inspection period after exposure for 2 years and for 4 years. In most soils, especially in the acid soils, there was some softening of the surface of the specimens. However, scratching several specimens indicated that this softening did not extend to greater depths than about one-thirty-second inch below the surface. The softening probably occurred only on the outer layers of the asbestos-cement sheet that were applied to the pipe without pressure during the manufacturing process in order to facilitate the removal of the pipe from the press. The material immediately under the softened outer layers appeared to be of the same density as the rest of the specimen.

Table 63 .- Absorption of water by asbestos-cement pipe.

	Specimen	v	Vater absorpti	on—percentag	e gain in weigh	t
Soil No.	No.	1 day	2 days	3 days	4 days	14 days
	SPI	ECIMENS 2	YEARS UND	ERGROUND		
1 3 5 6	41 68 149 39 21		1.64 2.13 1.86 .86 3.08	1.87 2.74 2.35 .97 3.86		3.68 6.75 6.76 1.56 7.49
0	71 12 56 1 111		5.21 1.13 1.55 1.50 1.15	6.26 1.36 1.81 1.97 1.31		10.18 2.79 3.05 4.25 2.70
5. 6. 7. 9.	121 99 131 89 101		1.58 1 11 1.00 1.96 2.05	1.69 1.37 1.25 2.48 2.40		3.99 4.42 5.10
	SPI	ECIMENS 4	YEARS UND	ERGROUND		
53 	67 147 31 26 74	1.69 1.75 2.53	2.26 2.58 3.14		3.04 3.80 3.57	
51 22 33 44	13 57 9 119 128	1.37 2.45	1.54 3.77 1.02		1.80 5.22 1.18	
69	97 133 87 109	1.85 1.35 2.14	2.04 1.77 2.61		2.34 2.11 3.24	
		UNBURI	ED SPECIM	ENS		
	2 3		4.73 5.96	5.69 7.26		

The specimens of asbestos-cement pipe removed after exposure of 2 and of 4 years were stored and were not subjected to physical tests until two years after the removal of the longer-exposed specimens, that is, until the specimens had been in storage for 4 and for 2 years, respectively. Then these specimens, together with five specimens which were obtained from the manufacturer at the same time as the others but which had not been buried, were subjected to several tests to be described.

Whether they were affected by the long periods during which they were in storage cannot be determined. Before crushing and bursting tests were made the specimens were immersed in water for at least 48 hours.

## (a) WATER-ABSORPTION TEST

Weighed air-dried specimens were immersed in water at room temperature for 2 to 14 days, after which they were removed and wiped with a damp cloth and reweighed. Water absorption was expressed as the percentage gain in weight. The results are tabulated in table 63.

There is no evident consistency in the amount of water absorbed by the specimens as a group, or by the individual specimens buried in the same soil and removed during different periods. Figure 32 shows some of the representative curves obtained from the data. The curves

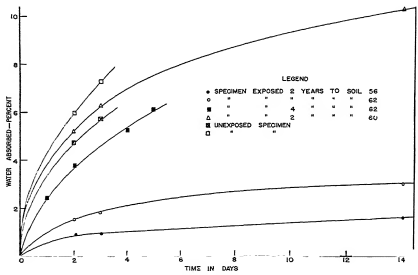


FIGURE 32.—Percentage of water absorbed by asbestos-cement specimens.

indicate that the unburied specimens absorbed more water than did the buried specimens, with the possible exception of the specimen buried 2 years in soil 60. The curves also indicate that the specimens were not saturated when tested.

## (b) CRUSHING TEST

The crushing tests were made on 5¾-inch lengths of pipe prepared by cutting in half each specimen that had been used for the water-absorption tests. After being air-dried these sections were immersed in water for 48 hours. The tests were made by using three-edge bearings according to the method described in the Federal specifications for asbestos-cement pipe. The load was applied at a uniform rate of approximately 1,000 pounds per minute until failure of the pipe occurred. In testing the specimens, the two sections from the same specimen were

placed under the hydraulic jack in such a way that the parts of the sections receiving the maximum stress were at an angle of 90 degrees to each other.

The crushing strength in pounds per linear foot for each of the sections was calculated. The results of these tests are given in table 64 for the specimens exposed to the soil and for the unexposed specimens. No evident correlation is shown between the values for the crushing strength and the age of the specimens either in individual soils or in any group of soils.

Table 64.—Results of crushing tests on the asbestos-cement specimens exposed to various soils and on the unexposed specimens.

		Specimens	s 2 years un	derground	Specimen	s 4 years un	derground
Soil No.	Sample	Minimum wall thick- ness	Crushing strength	Apparent specific gravity	Minimum wall thick- ness	Crushing strength	Apparent specific gravity
51	i b	\$\bar{v}_n\$.  0.76 .76 .72 .69 .69 .71 .71 .71 .72 .72 .73 .73 .73 .77 .77 .77 .77 .77 .72 .72 .72 .73 .73 .73 .73 .73 .73 .77 .77 .77 .77	lb per linear ft	1.90 1.90 1.90 1.96 1.86 1.94 1.889 1.889 1.889 1.880 1.890 1.891 1.893 1.992 1.995 1.994 1.888 1.992 1.995 1.994 1.888	in	lb per linear ft	1.92 1.83 1.86 1.884 1.91 1.90 1.99 1.99 1.99 1.99 1.99 1.99
		UNEX	POSED SP	ECIMENS			
4 5 Average of un-	( a b a b b				0.70 .70 .71 .71	9,940 10,270 11,840 10,530	1.92 1 90 1 90 1.93
exposed speci- mens					.70	10,640	1.91

## (c) BURSTING TEST

For the bursting tests the second of the pair of specimens removed from each test site at each inspection period was immersed in water for a minimum of 48 hours, and placed in the hydrostatic-pressure testing apparatus shown in figure 33. Internally fitting rubber cups were used to close the ends of the pipe. The apparatus was so designed that the pipe was not subjected to end compression during the test. After filling the pipe under test with water, the entrapped air was allowed to

escape, and the pressure was increased at an approximate rate of 10 pounds per square inch per second until the pipe failed. The pressure gage employed was calibrated before the series of tests.

Many samples failed by a small piece splitting out at one end with the extension of a crack from this point along the entire length of the specimen. The others failed by cracking in one or two places along the

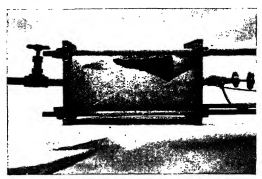


FIGURE 33 .- Bursting-test apparatus.

entire length of the specimen. In all but three cases, a crack went through a hole,  $^3\!\!/_6$  inch in diameter, near one end of the specimen, which was used to hold the identification tag. Here again there is no evident correlation between the bursting pressure and the age of the specimens either in individual soils or as any group of soils (see table 65).

Table 65.—Results of bursting tests on the asbestos-cement specimens exposed to the various soils and on the unexposed specimens.

	Specimens	s 2 years un	derground	Specimen	s 4 years un	derground
Soil No.	Minimum wall thick- ness along fracture	Bursting pressure	Apparent specific gravity	Minimum wall thick- ness along fracture	Bursting pressure	Apparent specific gravity
51	in. 0 71 .75 .72 .70 .73	lb/in. ² 995 1,140 1,085 995 1,140	1.89 1.83 1.87 2.00 1.76	in. 0.71 .72 .73 .71	lb/in.2 1,010 1,100 1,050 925	1.84 1.77 1.84 1.90
60	.71 .71 .73 .75 .67	1,010 1,205 1,095 1,175 1,100	1.76 1.92 1.97 1.84 1.98	.72 .71 .70 .74 .74	1,165 1,065 1,125 1,150 1,150	1.85 1.94 1.78 1.88 1.94
65. 66. 67. 69. 70.	.69 .75 .72 .74 .72	935 1,215 1,030 1,255 1,155	1.76 1.93 1.86 1.93 1.80	.70 .71 .73 .68 .77	1,070 1,240 1,105 1,215 1,285	1.84 1.89 1.82 1.89 1.84
	UNEX	POSED SP	ECIMENS			
1				0.71 .72 .73 .72	995 940 970 968	1.81 1.88 1.93

#### (d) APPARENT SPECIFIC GRAVITY

Samples from the specimens upon which the crushing and bursting tests had been made were dried in an oven at about 110° C for 18 hours. then cooled to room temperature, and the weight of the dry sample determined. The samples were then immersed in water for a period of 24 hours, and the weights in the wet condition were determined in air and then submerged in water. From the weights of the dry samples and the volume of water displaced, the apparent specific gravities of the specimens shown in tables 64 and 65 were calculated. With few exceptions, the specific gravities of the specimens fall between the values 1.80 and 2.00.

## (e) COMPARISON OF DATA FROM VARIOUS TESTS

Analysis of the data in tables 63 to 65 fails to bring out any correlation between any of the tests. The bursting- and crushing-test data for the specimens exposed for 2 and for 4 years do not show any evident differences. In some soils the 2-year specimens appear superior to the specimens exposed 4 years, and in other soils the reverse is true. No systematic differences between specimens exposed to different soil conditions can be detected.

Table 66.—Condition of parkway cable exposed 10 to 16 years.

G = good.	
$\mathbf{F} = \mathbf{fair}$ .	
B = bad.	

D = destroyed.

M = metal attack.
P = pitted.

[All steel is zinc-coated]

Soil No.	Duration of test	Outer fabric	Inner fabric	Outer steel	Inner steel	Lead sheath
1	Years 11.65 15.48 10.02 15.56 15.53	F  D G F	<del>ዕ</del> ውችዋው	R M P P	G M R SR M	G TW G TW TW
79	16.94 15.89 11.95 15.59 12.00	G B D	00000	P M P M P	SR R R M SR	W TW W TW
17	15 75 11.71 11.63 11.95 11.67	¥8444	00000	M R R G SR	M SR G G	TW TW TW G TW
26	15.90 9.60 10.08 17.04 15.73	F D B F B	GGFGG	M P P M P	SR P G SR M	TW TW TW TW TW
32	11.66 12 00 10.16 15.69 12.04	OFFFB	G G G	R SR P M SR	G SR G M SR	TW TW TW TW
38	15.82 12.00 17.41 12.02 11.73	FF BBB	G G F	PH SR P R R	R G SR SR R	TW TW TW G G
46 47	12.00 17.43	D G	G G	R M	R SR	TW W

However, the data do indicate that the asbestos-cement pipes generally gained strength during exposure to the soil, and that the softening observed on the outer layers of the specimens did not penetrate deeply. For only two specimens are the values for the bursting pressure less than the maximum value of the unexposed specimens, and in only one soil are both values of the crushing strength for the exposed specimens less than the average value of the unexposed specimens. It is probable that during exposure of asbestos-cement pipe to the soil, a curing process takes place which tends to increase the strength of the pipe, but there is no indication that the pipe is any stronger after 4 years of exposure than after 2 years of exposure.

## 5. PARKWAY CABLE

Parkway cable consists of one or more conductors so insulated and protected that the cable can be laid in a trench without further protection against moisture or mechanical injury. The cable tested consisted of a single rubber covered copper conductor with a lead sheath. Over the sheath was a wrapping of fiber treated to prevent rotting. This was surrounded by two spiral wrappings of zinc-coated steel tape. Over the tape was a wrapping of jute or similar material impregnated with a bitumen. When bent sharply the strands of fiber would separate. Table 66 shows the condition of the several parts of the cable buried for from 10 to 17 years. In no case was the usefulness of the cable impaired.

## VIII. RESULTS OF FIELD TESTS ON PROTECTIVE COATINGS

## 1. METALLIC COATINGS

(a) ZINC

As was shown in an earlier section, zinc corrodes under many soil conditions. It is a sacrificial coating first recommended for protecting iron by Sir Humphrey Davey [65] in 1825.

Zinc applied by the hot-dip methods is very extensively used for the protection of small-diameter underground pipes, especially those used for water services. The coating consists of an outer layer of nearly pure zinc and one or more layers of zinc-iron alloy, all of which are anodic to iron and steel and tend to reduce the rate of corrosion of the base metal if small areas are exposed.

The National Bureau of Standards included galvanized pipe in its 1924 burials, and 2 years later undertook to determine whether or not the composition of the iron to which the zinc was applied affected its protective value. The results of the tests indicated that any effect of the differences in these base materials was too small to be shown by the test methods employed, and from this it may be inferred that they are too small to be of any importance under service conditions.

An attempt was also made to determine whether or not the rate of corrosion of galvanized specimens was affected by the thickness of the alloy layer by testing coatings of approximately the same weight applied to No. 16 and No. 18 U. S. Standard Gage Bessemer sheet in 5 soils. Because of the uncontrolled variables involved or because such differences in the thickness of the alloy layer as were produced did not greatly affect the protective value of the coating, no difference in the protective value of the two coatings was found. The determination of the effect of the thickness of the alloy layer requires a closer control of

the factors affecting corrosion than can be obtained under normal field conditions and it can best be solved by laboratory experiments.

Table 67 recalculated from Research Paper RP982 [66] shows the corrosion of galvanized pipe and sheet exposed for approximately 10 vears to 43 soils. The losses have been recomputed as total losses per square foot, as a rate implies that the corrosion is proportional to the time of exposure which is not true of coated metals exposed to soil. To show the effect of the coating, the ratio of the average loss of the galvanized sheets to the loss of the unprotected pipe has been computed. Of course this ratio is different for different soils and would change with the thickness of the coating and the period of exposure.

Figure 34, devised by Ewing, presents the data graphically. The length of the heavy lines or horizontal columns shows the weight of the coatings plotted on the same scale as the loss of weight. Along these columns and their extensions the condition of each specimen has been correlated with the loss of weight of the specimen.

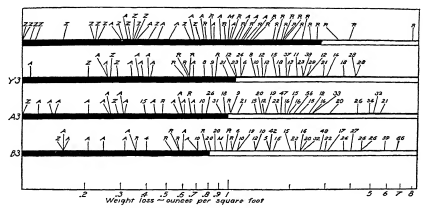


FIGURE 34.—Correlation between losses of weight and condition of galvanized specimens as determined by inspection.

The specimens are 10 years old. Where figures appear, they are the rates of penetration of the maximum pit, in mils per year. Where letters appear, they have the following meanings:

Z = Zinc continuous over specimen.

A = Blue or black alloy layer exposed over at least part of specimen.

R = Rusted or bare steel exposed.

M = Shallow metal attack; no pit as great as 10 mils total depth.

The heavy black lines indicate the thickness of the coating on the same scale as for the loss of weight.

It will be noted that with few exceptions, which may possibly be accounted for by thin spots in the coatings, the loss of weight at which rusting appears corresponds closely to the weight of the coating. This may be interpreted as indicating that corrosion of the base metal did not begin until nearly all of the zinc and zinc-iron alloy had been destroyed. The figures along the columns are maximum pit depths. They start near the ends of the solid columns. It will be noted that the heavy double coating prevented all pitting during the 10 years of the test but that rusting had begun in several soils. This indicates that the protective value of this coating is on the average approximately 10 years. The data do not show whether the rate of penetration of galvanized specimens that have begun to rust is as high as for ungalvanized specimens.

TABLE 67.---Corrosion of galvanized pipe and 16-gage steet sheet with nominal 2-ounce coalings [The numbers in parentheses are the weights (oz/ft*) of the contings.]

preimen.
c szposed over at least a portion of specimen.
sposed.
no pits as great as 10 mils- total depth.

The second secon			Кабен о	Rates of loss of weight (02/ft?)	eight (oz.	(£1J/	i		Condition	Condition or maximum penetration (in mils)	mum penc	trution (ii	ı mils)
VDe	Dura-	Pipe		<u> </u>	Sheet			Ratio	Pipe ^d ,		Sheet	ot	
	of test	A (2.82) ⁿ	A3 (0.99)	B (0.81)	Y3 (1.07)	Аverage	Barch		B (1.81)	₹	A3	<b>x</b>	X3
	Years 10 66 9.92 10.09 10.62	2.92 0.35 0.35 1.94 1.82	4.62 0.44 1.85 2.50	4.79 0.58 1.72 3.03	4.46 0.29 62 2.05 2.45	4.62 0.44 0.44 1.89 2.66	10.20	0.45	EEEE	HMMHH	28 23 12	28 R 10 17 22	30 14 12
r loam	10.16 10.48 10.63 9.48	0.12 2.62 0.78 1.10	0.27 2.69 0.65 .87 1.74	0.25 2.36 0.56 .78 1.71	0.00 3.04 0.39 1.08 1.75	0.20 2.70 0.53 1.73	5.55	190	- EEEE	RPZEZ	A 17 29 16	A 17 19 24	1238 22 1238 22
'm ly loam.	10.55 10.17 10.16 10.64 10.06	0.90 7 .87 26	0.42 .33 2.23 0.33	0.91 .96 .37 .34	1.15 0.26 1.37 0 68	0.83 .52 1.32 0.45	1.79		4 4	A RAZ	16 A Z Z	PPRM2	ZZ&*
	10.04 10.57 10.51 10.67 9.93	.99 3.64 0.68 1.22 1.19	1.46 5.90 0.82 2.10 2.04	2.04 3.92 1.16 2.40 2.28	1.78 3.89 1.04 1.56 2.37	1.76 4 57 0.67 2.02 2.23	6.44 3.30 5.01 7.16		RAR	MRZRR	20 22 27 17 16	15 27 20 21 32	18 19 22 13 29
ly loam	10.16 10.63 10.65 10.48 10.08	10.26 10.26 .36 .71	3.64 0.13 1.09 0.78 1.49	2.64 0.17 1.03 1.09 1.37	6.38 0.04 .92 .80	4.22 0.11 1.01 0.89	25.66	16	Z	822244	37 A 10 10 12	21 R 10	41 R 8 9

1.00   .01   .02   .72   .05   3.04   .21   R   R   R   A   A   A   A   A   A   A	-	5.19 1.58 0.05	4.48 1.42 0.17	4.24 1.48 0.05	4.0 1.45 1.64 1.64	14.70	18:	30		888	 882.	88 I
10   16   0.38   0.770   1.07   0.443   0.752   11.30   .22     R		19. %	.62	.72	.05	3.04			4 4 4	24	AH.	84
10.02   f_2.03   0.15   0.20   0.12   0.17   0.17   0.25   0.12   0.17   0.15   0.17   0.15   0.17   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15   0.15	10.16	0.70	1.07	0.43	30.73	11.90	77	4	34Z	4 A A 1	88.4 4	74 A
10.05	10.62 (0.21	0.16	0.22	0.12	3.08 0.17	8 54	36		22	A 20	17 A	414 V
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	10.05	0.22 22.88	0.38	0.27	0.31	7.48	29	ZZ	R A	47 A	48 A	38 A
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	10.73 1.38	0.22	6.27	.69	2.68	12.72	.21	25	¥4 :	PR2	63 A	% ₽¤3
1.00 1.00 A A B	10.55 f 1.84 10.54 0.17 10.60 1.06	1.01 0 111 .70	1.69 0.17 .83	1.79 0.28 .69	1.50 0.19	13.53	.04	88 A A 38	ZZ	19 Z	Z ^O ZP	16 Z

an here is in ounces per square 1000 to express of rolled iron and steel specimens puried a summa rough.

In the presents the ratio of the average of the sheets to the bare-metal data column, a low figure indicating that the galvanizing was effective in reducing of the ratio of the average of the sheets to the bare-metal data column, a low figure indicating that the galvanizing was effective in reducing the ratio of the average of the sheets to the bare-metal data column, a low figure indicating that the galvanizing was effective in reducing the ratio of the average of the sheets to the bare-metal data column, a low figure indicating that the galvanizing was effective in reducing the ratio of the average of the sheets to the bare-metal data column, a low figure indicating that the galvanizing was effective in reducing the ratio of the average of the sheets to the bare-metal data column, a low figure indicating that the galvanizing was effective in reducing the sheets to the sheets to the bare-metal data column, a low figure indicating that the galvanizing was effective in reducing the sheets to the sheets to the bare-metal data column, a low figure indicating that the galvanizing was effective in reducing the sheets to the sheet 12 years. They were not weighed before burial so weight losses are not known, pecimens. The condition or penetration is for the worse-corroded specimen, I this material. The condition is for the worse of these specimens. Figure 35 from the report referred to shows in a simpler way the relation of pit depths to loss of weight and in addition correlates these data with the test-site numbers. The ratio of the pit depth to the loss of weight represented by the slope of the curve is proportional to the

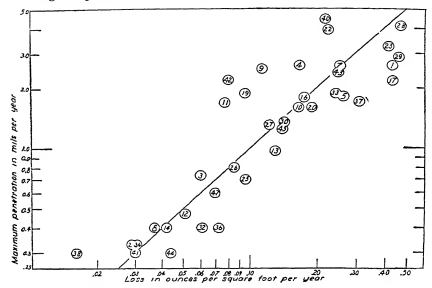


FIGURE 35.—Correlation between rates of loss of weight and rates of penetration on galvanized specimens.

pitting factor. As for the unprotected steel, this appears to differ with the soil. In the case of the data represented in figure 35 the dispersion of the data may be affected by variations in the thickness of the galvanizing, which according to Rawdon, [67] on a sheet having an area of 24 square feet and an average coating thickness 2.67 ounces per square foot may have in places as little as 1.79 ounces per square foot or as much as 3.06 ounces per square foot over an area of 18 square inches. Table 68 shows the loss of weight and maximum penetration of galvanized iron pipe carrying a heavier coating exposed to 14 soils for 4 years. A few of the test sites listed in tables 67 and 68 are identical although they differ in identification numbers. The only known difference in the coatings that may account for the much better performance of the older one is that it was put through the zinc bath twice.

Ewing made an extensive study of the corrosion of gas service pipes, and concluded that galvanized coatings appear to have little value in reducing pipe replacements in congested cities and should not be recommended as a cure for corrosion of services if it is at all severe. He attributes the discrepancy between the performance of the older specimens in the Bureau tests and the performance of galvanized service pipes to electrolysis and galvanic action between the galvanized pipes and ungalvanized mains to which they were connected. These factors are undoubtedly important. It is possible, however, that the galvanized

specimens in table 68 more nearly represent the galvanized material in service. Ewing presented no data on the weight of zinc on the services on which he reported. He suggests that electrolysis and galvanic action might be reduced by insulating the service from the mains. If this is done, a long leakage path between the main and the service should be provided to avoid concentrated corrosion near the insulating point.

Table 68 .- Loss of weight and depth of maximum penetration of galvanized and black iron pipe exposed for 4 years.

	Soil	Dura-	Galvani 3.08	zed pipe oz/ft²		k iron pe	Condition
No.	Type	tion of expo- sure	Loss of weight	Penetra- tion	Loss of weight	Penetra- tion	of coatinga
53 55 56 58	Cecil clay loam. Hagerstown loam. Lake Charles clay. Muck. Rifle peat	Years 4.01 3.90 3.99 4.01 3.98	oz/ft ² 1.39 b.1.22 3.89 5.40 7.18	Mils 6 8 7 b 21 12	oz/ft ² 2.86 2.60 16.03 8.78 8.06	Mils 98 50 104 46 5 38	2 2 2 3 3
61 62 63 64 65	Sharkey clay	4 01 4 00 4.01 3.98 3 99	1.46 2.28 2.15 1.58 b.2.25	12 9 10 9 6	4.99 4.30 9.20 5.96 4.56	45 56 38 67 59	2 2 1 2 3
66 67 69 70	Mohave fine gravelly loam Cinders	3 95 3.98 3 98 3.98	3.32 5 40 3.37 4.52	8 45 11 12	12.31 4 37.03 3.28 5 9.72	° 145+ 145+ 20 118+	23333

a 1 = coating on more than 50% of surface.
2 = coating on less than 50% of surface.
3 = little or no coating remaining.
b Data for individual specimens differed from each other by more than 50%.

• +=Hole in 1 or both specimens due to corrosion.

d Data for 1 specimen; the other specimen was destroyed.

The extent to which zinc applied to iron protects the iron after it has been exposed by abrasion or corrosion has been studied by several investigators. Mills [68] reported that by casting a zinc collar 4 feet long around 20-foot sections of oil-well tubing the Empire Gas & Fuel Co. extended the life of the tubing from 2 weeks to more than 8 months. Whether this is analogous to the protection afforded by a zinc coating may be questioned.

The first National Bureau of Standards soil-corrosion report [69] describes an experiment in which the zinc was machined off from a 17-inch length of 2-inch galvanized iron pipe to within 2 inches of each end. The pipe was immersed in tap water. Rust formed on the pipe to within 2 inches of the galvanized surface. It has been suggested that the protected lengths depends on the conductivity of the electrolyte.

The way in which a coating of zinc or an attached piece of zinc furnishes protection to iron has been explained in several ways. In some electrolytes a film of corrosion products may afford most of the protection. This is particularly applicable to coatings of zinc. Protection to iron may be afforded by the action of zinc as a sacrificial anode. Bannister [70] has suggested that the protection afforded by zinc plates in a boiler is due to the absorption of oxygen by the zinc. Under favorable conditions zinc coatings could act similarly. Probably each explanation accounts for the protective effect of zinc under some conditions but does not account for it under others.

### (b) LEAD

Lead-coated pipe specimens were buried in 41 soils in 1923-24 and in 15 soils in 1932. The maximum and minimum thickness of lead on the 1923 specimens were 2.50 and 0.3 mils, respectively; the corresponding values for the 1932 specimens were 2.97 and 0.5 mils. The last of the 1923 specimens were removed from the less corrosive soils in 1941. Table 69 shows comparable data for these coated specimens and for uncoated steel pipe exposed for somewhat longer periods. Table 70 gives similar data for the same material exposed to more corrosive soils. In the latter table, corrosion is expressed in terms of rates because the steel specimens were exposed for somewhat longer periods. As rates change with the period of exposure, this is not an entirely satisfactory way to report corrosion. However, the table is useful for the comparison of the relative merits of coated and uncoated steel after an exposure of approximately 10 years.

Table 69.—Loss of weight and depth of maximum penetration of 11/2-inch leadcoated pipe and Bessemer steel pipe exposed 16 years."

	Soil	Lead-	coated	Bessem	er steel	Condition
No.	Type	Loss of weight	Penetra- tion	Loss of weight	Penetra- tion	of coating
5 6 7 9	Dublin clay adobe. Everett gravelly sandy loam. Maddox silt loam. Genesee silt loam. Hanford fine sandy loam.	.36 4.42	Mils 104 14 145+ 600 d 28	oz/ft ² b 7 . 12 2 . 03 5 . 82 5 . 80 6 . 04	Mils 50 18 50 64 72	3 0 3 2 2
17 24 26 27 31	Keyport loam Merrimac gravelly sandy loam Miami silt loam Miller clay Norfolk sand	1.49	49 28 48 47 28	9.54 1.79 4.28 10.14 3.72	41 16 42 69 43	3 2 2 3 1
35 36 38 41	Ramona loam Ruston sandy loam Sassafras gravelly sandy loam Summit silt loam	b .14 1 27 2 .17 1 .27	ь 10 24 34 80	1.51 4 13 2.30 6.87	8 55 28 92	0 3 3 2

Lead-coated pipe buried in 1923-24.
 Bessemer steel pipe buried in 1922.
 Data on 1 specimen only.
 Average of 3 specimens.
 Average of 4 specimens.

^{• 0 =} coating over entire specimen.

^{1 =} coating on more than 50 percent of surface. 2 = coating on less than 50 percent of surface. 3 = little or no coating left.

Table 70.—Loss of weight and depth of maximum penetration of 11/2-inch lead-coated pipe and Bessemer steel pipe exposed for 10 years.

			- P-PO OIO		1 10 gea		
		Rates	of loss (oz	/ft²)/yr		of maxim	um pene- s/yr)
Seil	Soil type	Lead- coated steel	Bare steel a	Lead cable sheath a	Lead- coated steel	Bare steel	Lead cable sheath
1 2 3 4 5	Allis silt loam. Bell clay Cecil clay loam Chester loam Dublin clay adobe.	0.529 .102 .066 239 .597	0 80 .49 .43 .52 .45	0.173 .067 .059 .177 .135	11.9 4.0 3.6 7.0 6.9	5.8 6.7 4.5 6.8 3.1	8.1 1.8 1.4 4.0 3.8
6 7 9 10 11	Everett gravelly, sandy loam Maddox silt loam Genesee silt loam Gloucester sandy loam. Hagerstown loam	.096 .207 .080 .163 .071	.08 .37 .43 .36 .16	.025 	2.0 7.0 5 0 6.5 3.4	1.4 2.5 4.5 3.2 4.9	0.8  1.2 1.2
13 14 15 17 18	Hanford very fine sandy loam	.064 .036 .052 .379 .046	.39 .65 .77 .23	.049 .036 .028 .016	5.4 4.2 3.2 4.7 4.6	7.1 4.4 3.3 3.2	1.7 2.2 1.7 1.1
20 22 24 27 28	Mahoning silt loam Memphis silt loam Merrimac gravelly, sandy loam Miller clay Montezuma clay adobe	.224 .192 .033 .231 .314	.52 .61 .12 .63 1.75	.268 .085 .015 .067 .069	5.3 7.3 2.2 5.6 7.8	5.5 6.6 2.1 4.9 15.1	4.1 1.2 1.5 2.4 0.9
29 31 32 33 35	Muck Norfolk sand Ontario loam Peat Ramona loam	.689 .028 .072 .537 .011	1.61 0.22 .32 1 22 0.09	.343 .022 .028	6.3 1.5 5.2 7.7 (b)	12.0 2.5 4.3 8.9 0.4	.9 .5 1.5
36 37 38 40 41	Ruston sandy loam St. Johns fine sand Sassafras gravelly, sandy loam Sharkey clay Sumnit silt loam	.032 .202 .053 .191 .051	.24 .58 .22 .56 .45	.032	1.8 6 4 3 4 6.2 5.8	3.9 5.6 2.5 6.6 7.6	1.1
42 43 45 46 47	Susquehanna clay Tidal marsh Un'dentifed all'all soil Un'dentifed sar dy loana Un'dentifed sir loana	.092 .726 .198 .033 .184	.95 1.47 0.79 .37 .23	.019 .021 .015 .122	4.6 18 8 9.2 4.5 6.1	7.1 8.7 6.9 8.8 2.4	1.0 1.7 1.0 5.7

^{*} Bessemer steel pipe  $1\frac{1}{2}$  by 6 inches and commercial lead cable sheath in sheets  $20\frac{3}{4}$  by  $3\frac{1}{4}$  by 0.112 inch removed from sites at the same time the lead-coated specimens were removed, were buried, in most cases, 1 or 2 years longer. The figures given are the averages from 2 specimens in the case of the steel and 1 specimen in the case of the lead sheath. The penetration of the sheath was determined by averaging 2 pits, each of which was the deepest on 1 side of the specimen. These pits were, of course somewhat deeper than would be expected on specimens of the same exposed areas as that of the pipe.

b No pits.

Table 71 gives similar data for somewhat larger and more heavily coated specimens exposed to other corrosive soils for 9 years. Although, on the whole, the lead-coated specimens showed less penetration than the unprotected steel, their performance can be considered satisfactory in only a limited number of corrosive soils chiefly those containing sulfates. As the data on lead pipe (table 60) show that in a considerable number of soils the penetration is much deeper than the thickness of any coating of lead that has been applied commercially, it seems doubtful that lead coatings should be recommended except, perhaps, in the very limited number of cases where the soil is known not to attack lead.

TABLE 71Loss o					of	lead-coated	and
	stee	l pipe expo	sed for 9 ye	ears.			

	Soil	Lead	l-coated a	Low-e	arbon steel N	Condi- tion of
No.	Type	Loss of weight	Penetration	Loss of weight	Penetration	coat- ing b
53 55 56 58 59	Cecil clay loam Hagerstown loam Lake Charles clay Muck Carlisle muck	oz ft ² c 1 . 12 0 . 76 20 73 14 85 1 . 07	Mils c 41 44 d 145+(7) e 91 f 21	oz/ft ² 4.09 3.82 28.76 16.24 4.70	Mils 59 59 154+(7) 110 140	2 1 3 3 1
$60 \\ 61 \\ 62 \\ 63 \\ 64$	Rifle peat Sharkey clay Susquehanna clay Tidal marsh Docas clay	f 7.07 3 89 3.64 3.52 2.10	50 50 59 • 145 + 73	16.72 5.78 6 65 f 9.03 g D	ef 27 f 96 f 87 54 154+(5,7)	3 2 3 2 2 2
65 66 67	Chino silt loam.  Mohave fine gravelly loam Cinders	3.22 3.39 D	72 57 145+(5, 7)	12 86 18.56 h 58.39	$ \begin{array}{c c} 112 \\ 154 + (5,7) \\ 154 + (2,5,7) \end{array} $	2 2 3

a This coating was 0.00144 inch thick and contained 1 percent of tin.

b 1 = coating on more than 50% of surface. 2 = coating on less than 50% of surface.

#### (c) ALUMINUM (CALORIZED)

A calorized coating is an alloy of aluminum with the base metal [67]. It was developed primarily for the purpose of preventing scaling when the metal is subjected to high temperatures. The aluminum oxide, which develops on the surface of calorized materials, was considered as a possible protection against soil corrosion. There are two processes of calorizing, one known as the powder or dry process; the other as the dip, or wet, process. Specimens of pipe calorized by each process were placed in seven soils in 1924. Table 72 gives the rates of corrosion of the calorized specimens, together with the corresponding data for unprotected steel after 10 years of exposure.

In all cases the calorized specimens lost less weight and, with one exception, were pitted at a lower rate than the unprotected pipe in the same soils. In five of the six soils, the powder-calorized pipe was pitted at a lower maximum rate than the dip-calorized pipe, whereas in five of six soils the dip-calorized pipe lost less weight than the powder-calorized pipe. It is thus apparent that the powder-calorized coating was more effective in reducing pitting, although it allowed a greater loss of weight

than the dip-calorized coating.

The thickness of the coating on both types of calorized pipes was measured in several places, by the use of the chord method,6 potassium ferrocyanide and copper sulfate being applied to distinguish the alloy boundary. The thickness of the coating on both kinds of calorized pipe was found to be reasonably uniform. The thickness of the powdercalorized coating was 2 mils and that of the dip-calorized was 11 mils.

It is evident from table 72 that although, on the whole, calorizing made the pipe more resistant to soil corrosion, the process as applied to

^{2 =} coating on less than 50% of surface.
3 = little or no coating left.
c Data for only 1 specimen.
d +=1 or both specimens punctured. A number in parentheses after the pit depth indicates that 1 of both specimens were punctured in previous removals, i.e., (5) indicates a puncture after 5 years, etc.
c Uniform corresion—no reference surface left on pipe.
f Data for it illustry specimens, cultivate room the average by more than 50%.
b D=both specimens destroyed by corrosion.
h Data for 1 specimen only; the other specimen was destroyed.

⁶ See reference 67 for a description of calorizing processes and the chord method.

Table 72.—Corrosion of calorized steel pipe exposed for approximately 10 years.

Soil	Soil type	Rates	of loss of (oz/ft 2/yr)	weight a	Rates of	maximum n ª (mils/y	penetra- r)
		Dry calorized	Wet calorized	Bare steel b	Dry calorized	Wet calorized	Bare steel b
13 24 28 29 42 43 45	Hanford very fine sandy loam. Merrimac gravelly, sandy loam. Montezuma clay adobe. Muck. Susquehanna clay. Tidal march. Wyo.)	0 137 .019 .437 .250 .452 .861 .712	0.087 .013 .458 .088 .434 .184	1.24 0.12 1.75 1.39 1.25 1.72 1.23	3 4 2.4 5.5 3.4 7.0 3.1 3.1	4.4 3.2  7.8 4.2 3.4 4.4	13.0 2.1 15.1 9.2 9.2 7.4 11.9

a Average of 2 specimens.

Brates for unprotected Bessenier steel specimens of similar area which were exposed, except in the case of soil 13 for a similar period. The data presented for bare steel in soil 13 are for specimens exposed only 6 years, there being no data available for a longer period.

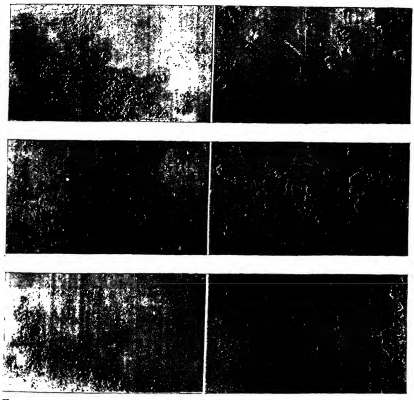


Figure 36.—Panoramic pictures of 2-in. calorized pipe exposed approximately 10 years.

The specimens on the left are dry calorized; those on the right are wet calorized. The soils to which the specimens were exposed were: top, Hanford very fine sandy loam; center, Muck; bottom, Susquehanna clay.

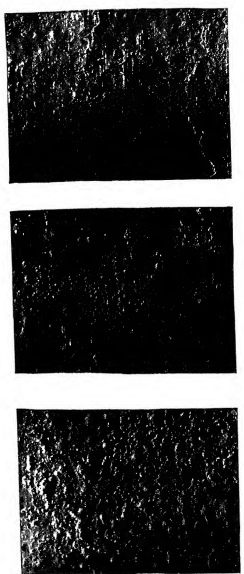


FIGURE 37.—Panoramic pictures of 2-in. steel pipe exposed with the calorized specimens shown in figure 36.

Top, Hanford very fine sandy loam; center, Muck; bottom, Susquehanna clay.

the specimens did not prevent pitting in any soil. The rate of pitting and loss of weight are usually high in poorly drained, corrosive soils where the use of a good coating would be economical. The appearance of the calorized specimens from three widely different soils is shown in figure 36. Unprotected steel specimens exposed to the same soils for the same times are shown in figure 37. While these photographs have been selected to show the appearance of more or less typical specimens, they are not presented as the basis for general conclusions.

### (d) TINNED COPPER

Table 73 shows the loss of weight and maximum penetration of tinnedcopper tube exposed for 4 years and similar data on deoxidized copper pipe exposed at the same test sites for 5 other years. The table shows that, in general, the coating of tin temporarily reduced the rate of loss of weight but, in some cases at least, apparently accelerated the rate of maximum penetration. The coating of tin was quite thin and much of it had disappeared from the specimens from most of the soils.

Table 73 .- Loss of weight and depth of maximum penetration of tinned-copper tubes exposed for 4 years and copper pipe exposed for 5 years.

M, shallow metal attack as indicated by roughening of the surface. P, definite pitting but no pits greater than 6 mils. +, one or both specimens contained holes due to corrosion.

	Soil	Tir	ned-coppe	r tube	Deo	xidized cop	per pipe	<u> </u>
No.	Туре	Expo-	Loss of weight	Maximum penetra- tion	Expo- sure	Loss of weight	Maximum penetra- tion	Condi- tion of coat- ing a
53 55 56 58 60	Cecil clay loam Hagerstown loam. Lake Charles clay. Muck. Rifle peat.	3.90	oz/ft ² 0.086 .11 .36 .64 4.54	Mils 7 M 11 39 42	Years 5.46 5.20 5.44 5.50 5.25	oz/ft ² 0.15 .14 .51 1.56 3.82	Mils 7 P P 12 28	1 0 2 2 2 3
61 62 63 64 65	Sharkey clay Susquehanna clay Tidal marsh Docas clay Chino silt loam	4.01	0.32 .081 2.19 0.22 .12	18 P b 14 6 P	5.50 5.47 5.55 5.22 5.26	0.35 .26 2.45 2.22 1.02	23 9 M 9 10	2 1 3 2
66 67 69 70	Mohave fine gravelly loam. Cinders	3.95 3.98 3.98 3.98	.23 20.37 ° 0.23 .074	P 60+ P 6	5.28 5.26	0.75 9.33	7 54	2 3 1 1

Aside from the question of minimum thickness required for protection, a possible cause of failure of tinned copper in soils is the reversal of potential of this couple. As tin is probably anodic to copper under all conditions, corrosion of tin would be expected to protect copper cathodically in the same manner that the corrosion of zinc protects the underlying steel in galvanzied materials. The reversal of potential may be due to the formation of tin-copper alloys, which have been shown under certain conditions to be more cathodic than copper [71].

a 0 = coating present over entire surface.

1 = coating present on more than 50% of surface.
2 = coating present on less than 50% of surface.
3 = little or no coating left.
b Uniform corrosion—no reference surface left.
c Data for 1 specimen only.

# 2. BITUMINOUS COATINGS

# (a) SCOPE AND METHODS

The National Bureau of Standards began testing pipe coatings as a means of preventing electrolysis about 1912 [72] and included several pipe coatings in its soil-corrosion tests in 1922. Each time specimens were buried additional samples of coatings were included. The tests indicated that as a means of preventing electrolysis, coatings were unsatisfactory, since they concentrated the current discharged when they failed, as most of those tested did within a short time. In the early tests only one bituminous coating, a cotton fabric impregnated with coal-tar pitch, appeared to be reasonably satisfactory, and the later development of better coatings makes superfluous any detailed report on the early Bureau tests. The Bureau's soil-corrosion investigation and corrosion losses of pipe-line operators led the American Gas Association and the American Petroleum Institute to employ Research Associates to cooperate with the National Bureau of Standards in field tests of pipe coatings. Field tests of coatings were started by the American Gas Association in 1929 and by the American Petroleum Institute in 1930. In general, the plan of testing consisted of applying coatings to sections of pipe or pipe lines located in selected soils and examining the coatings from time to time to determine the effectiveness of the protection which the coatings afforded to the pipe. Four criteria of the performance of the coating were used: 1. Appearance of the coating with respect to adhesion, cracking, distortion, and other physical changes; 2, the pattern test; 3, the conductance or resistance of the coating; and 4, the extent of the corrosion of the protected pipe.

Appearance: An association of coating manufacturers drew up an extensive code [29] for the visual inspection of coatings. This code was followed by both of the Research Associates. The code was intended to yield detailed information free from any bias of the inspector. The amount of data called for this code is too voluminous for presentation to the general reader. The pattern and conductance tests were developed by Ewing [29], Scott [73], and Shepard [74].

Pattern test: The section of coating is washed and painted with a slurry of kaolin or iron-free clay to fill the holes, cracks, and depressions. Over the clay is placed a wet sheet of heavy absorbent paper, such as that used by plate printers as wiping paper. One manufacturer sells it for coating tests as "pinhole papers." Any tough, thick paper that will absorb a considerable quantity of water quickly would be satisfactory. The paper is usually cut in lengths equal to the circumference of the pipe and I foot wide. Its position on the pipe is marked, so that if desired, it can be replaced in its original position. Over the paper is wrapped several layers of wet cotton flannel or outing cloth, which is held in place by means of a saddle made of several strips of sheet copper held together at the ends by cross strips. A battery and a voltmeter are connected between the saddle and the pipe for such a time that the product of the voltage multiplied by the time in minutes is 100. If patterns are to be compared, this product must be kept constant. The pipe must be the anode. The paper is then removed from the pipe and washed in a solution consisting of 7 grams of potassium ferricyanide per liter of water. (This is approximately equivalent to 2 tablespoonfuls per gallon of water.) If current has flowed from the pipe through holes in the coating, ferrous hydroxide will be deposited in the paper and this will be turned blue by the solution. The result is one or more blue spots

showing the location and approximate size of the holes in the coating. The pattern serves as a record of the test and assists in the location of

the pinholes if they are small.

Conductance test: If the current and voltage are observed during the above test, the resistance or conductance can be computed, provided care is taken to avoid leakage by keeping the coating dry on either side of the pad. After these tests are completed, the coating is removed and the condition of the pipe is determined.

# (b) AMERICAN GAS ASSOCIATION TESTS

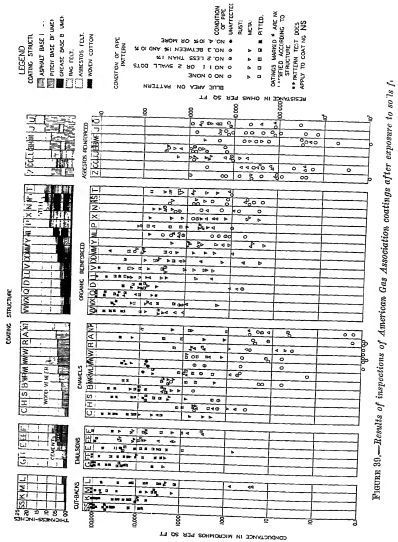
The American Gas Association, independent coating manufacturers, and the National Bureau of Standards undertook field tests of coatings applied to short lengths of pipe. Forty-two varieties of proprietary bituminous coatings described in table 17 were applied by the makers of the coatings to 20-foot lengths of sandblasted 2-inch steel pipe at a central coating plant. The pipe was then cut into 2-foot lengths, and the ends were closed by tin cans filled with the coating bitumen. The cans served



FIGURE 38 .- American Gas Association coatings before burial.

to keep water out of the pipe, protect the cut ends of the coatings, and support the specimen during shipment. Sixty samples of each material were provided so that four could be placed in each of fifteen soils. The coatings were removed and examined from time to time. Three inspections were made, the last one when the coatings had been exposed for about 4 years.

Figure 38 shows the specimens at one test site before burial. Figure 39, devised by Ewing [75], is a nearly complete report of the results of



his last inspection of these coatings. One set of coatings was still in the ground in 1944. As a result of his tests, Ewing concluded (1) that none of the coatings entirely prevented corrosion at all test sites, (2)

that coal-tar-base materials are more stable and waterproof than asphalt-base materials, but that a good coating could be made from either material, (3) that machine-applied coatings were superior to similar hand-applied coatings and advised against hand-application in the field, and (4) that any organic reinforcement in a coating is a weakness, especially if the bitumen is asphalt. He further advised against the use of any coating that would not afford some protection in poorly drained soils.

## (c) AMERICAN PETROLEUM INSTITUTE TESTS

(1) Sites and Coatings.—The tests conducted by Ewing were primarily demonstrations of the relative usefulness of certain proprietary coatings. The American Petroleum Institute undertook in 1930 a series of coating tests intended primarily to establish the engineering principles that should govern the design and selection of pipe-line coatings. Substantially the same manufacturers cooperated as in the American Gas Association tests. In addition to the establishment of the engineering principles that should govern the manufacture, selection, and application of pipe coatings, the planners of the Institute tests hoped to determine the applicability of tests of coatings, such as those made by Ewing, to the performance of coatings on pipe lines, and the applicability of the results of corrosion tests, such as the National Bureau of Standards was conducting, to the determination of corrosion of pipe lines.

The design of a series of tests that would accomplish these purposes was a complicated problem, involving many variables and requiring many compromises among the planners, and there was some uncertainty as to the significance of the resulting data. A somewhat detailed description of the manner in which the tests were carried out is, therefore, necessary.

Arrangements were made with 16 pipe-line operators to furnish 1,000 feet of new or substantially new operating line, together with labor for taking up the pipe, applying the coatings, and relaying newly coated pipe. Usually, sections of line where corrosion had occurred were selected, but accessibility of the line and the characteristics of the soil were also considered. In two cases the chosen section of the line had been in the ground a few years and was slightly pitted. In two other cases the soil was so corrosive that the line owners did not wish to take a chance on leaks and therefore laid new pipes parallel to the operating ones. Parts of these new lines were connected electrically to the pipe lines but did not carry oil. One line carried hot oil.

After the pipe had been carefully cleaned, a coating was applied to 30 feet of the line by or under the supervision of the manufacturer of the coating. Another manufacturer applied his coating to the next 30 feet of the line. The next 20 feet of the line was left uncoated as a control, and then two other coatings were applied. The process was continued until half of the new pipe had been used. The other half was then treated similarly. Thus at each site there were two 30-foot sections of pipe line coated with representative samples of each coating, and at one end or the other of each coated section there was a section of bare pipe.

A record was made of the resistivity of the soil opposite each coating to show the uniformity of soil conditions. The thickness of each coating was measured at a number of places to determine its uniformity. Before the coated pipe was returned to the trench, the coatings were carefully inspected and all discovered imperfections were repaired. To avoid injuries to the coating, great care was exercised in returning the pipe to the trench and in backfilling the trench. The location of the test sites, the soil types, and the profile descriptions are given in Appendix 2.

Tables 14, 15, and 16 give the descriptions of the coatings on the short pipe sections and on the line pipe and their characteristics. It has been suggested by Putnam [76] that alternating bare and coated sections of the line set up a condition of differential aeration that tended to accelerate corrosion under the coatings and to make the uncoated sections cathodic. Such a condition would tend to make the effectiveness of the coatings, as indicated by the condition of the pipe, appear less than it would have been had the entire lines been coated with one material. On the other hand, any stray currents in the earth would be collected and discharged chiefly by the bare sections, and thus the coatings may have been subjected to less than normal electrical stress. Similar arguments might be applied to the comparison of the corrosion of small uncoated pipe with that of the bare sections. However, a careful examination of the data does not disclose any definite evidence to support any of the above criticisms of the tests. This in part may be due to the fact that soil-corrosion data are always more or less variable. Small earth currents flowed in different directions at different times and places, and the soil conditions, as shown by earth resistivity measurements and by visual inspection, were not entirely uniform. It seems probable that, on the whole, the tests were as well conducted as they could have been, and it is doubtful that another test planned in the light of the old one would produce better results.

To determine whether or not the coatings applied to the lines were typical of their classes, and to get some idea of the relative merits of the various available coatings of each class, as well as to compare the results of tests of coatings on operating and on isolated short sections of small-diameter pipe, all the line coatings and all other bituminous coatings offered for test were applied to 2-foot sections of 3-inch boiler type as in the American Gas Association tests, and buried in a separate trench parallel to the operating line. Three specimens of each coating were buried at each test site. The characteristics of these coatings are shown in table 16. To supplement the bare control sections of the pipe lines, and to provide an additional measure of the uniformity of the soil and additional controls, as well as to study the relative rates of corrosion of working lines and small sections of pipe, several 2-foot sections of 3-inch bare steel pipe were buried in the pipe-line trench opposite the bare and coated sections of the line. The condition of the coatings was determined by the methods used in the American Gas Association tests. In the final inspection, the line pipe was marked off into 1-foot lengths, and the depths of the five deepest pits on each section were measured. The American Petroleum Institute tests were under the direct supervision of Scott [73] who reported on the first three inspections of the coatings. The final report on the tests was prepared by K. H. Logan [61] with the advice and assistance of a number of men representing pipe-line operators and coating manufac-

turers. This is by far the most extensive test of coatings ever conducted. The report presents a large quantity of data, which permit comparing the coatings in a number of ways. Data are also given regarding the uniformity of test conditions and to enable one to judge the relative

merits of different criteria of coating performance. For the present purpose, however, it is sufficient to present data that will permit a comparison of the coatings on the basis of the condition of the pipe which the coatings were designed to protect. The following section is

made up largely of abstracts from the final report.

(2) Coatings Applied to Pipe Lines.—Table 74 shows for each line coating at each test site, the average of the deepest pit on each of approximately 20 one-foot sections of coated pipe, the average of the deepest pits on 14 one-foot sections of adjacent uncoated pipe, and the deepest pit on the 3-inch control pipes buried in the same trench and having approximately the same area as 1 foot of line pipe. As an indication of the uniformity of soil conditions, the resistivity of the soil opposite the coated and uncoated sections of the line is also given.

In many cases the same basic coating material was used in two ways, for example, with and without a reinforcement or shield. Table 74 indicates which coatings are to be compared to determine the effect of the reinforcement. For a more complete description of the coatings,

see tables 14 and 15.

TABLE 74.—Averages of maximum pit depths on coated and

										-	Sites 3				1					
					Tem	I Temple, Tex,	×			Arl	сапкав	II Arkansas City, Kans.	Kans.			m m	III Beaumont,	III ont, Tex.		
		(		Pit depths (mils)	pths s)		Resis	Resistivity		Pit	Pit depths (mds)		Resi	Resistivity		Pit c	Pit depths		Resis	Resistivity
-	1		Line		Control	To.	(ohr.	rem)		Line	ပ်	Control	(g)	m-em)		Line	Cor	Control	(ohm	(ohm-cm)
Symbol Thickness (mils) Associated	Buitsoo	Character	Coated	Nearest bare	erisoqqO gairson	Opposite testest stad	Opposite guitsoo	estisoqqO tserserr erred	Deteco	Nearest bare	estisoqqO gaistee	Opposite testest exec	Opposite gaitsoo	etieoqqO teensen ensed	Coated	Nearest 5126	edieoqqO gailesoo	əfisoqqO fastasar ətad	etisoqqO gaitsoo	etisoqqO taeraen etaeraed
		1	COLL	APE	LICA	TIONS	3: CU.	COLD APPLICATIONS: CUTBACKS,	KS, C;	AND	EMU	AND EMULSIONS,	R, A							
C 21 Y A 65 F	75	Cutback coal tar	3.6	22 62	49	45 	490	525 525		.09	.19	43	1,200	875	208	185	HH	шш	590	480
							2. El	2. ENAMELS	LS											
N 69 N W 69 U W 88 H	75D :H	Coal-tar-asphalt enamel do Coal-tar enamel do	29	29		- 69	455	480	8 8 8 5 8 5 7	8888	48 39 49	22233	1,000 1,025 2,325 3,275	715 900 900	73 109 231 231	209 175 151 175	шшшш	HH .H	480 545 545 555	490 415 585 415
							3, 18	MASTIC	ا ت											
519		Asphalt mastic	1.5	21	- 89	61	200	700	700   0.25	48	52	46	1,800	1,025	0	151	H		360	585
- 1	- 1	4.	SHIELDED	COA.	ring;	S: CU	(BAC)	KS, C;	EMU	LSION	IS, A;	AND	COATINGS: CUTBACKS, C; EMULSIONS, A; AND ENAMELS	TELS						
29 C 419 A 63 N 81 L		Cutback asphalt	1.2	220	59	40.	525	495	1111	48:	::8	56	1,225	850	14 87	185	HH	HE	500	480
					70	6. REII	NFOR	REINFORCED COATINGS	CATI	NGS										
S 150 E 151 X 201 X 201 X 351 X 351	X   X	Grease Asphalt do do do do Coal-lar-usphalt enamel Coal-tar enamel	6.0 29 4.8 4.8 7.6	8888888	842 84 85 85 85 85 85 85 85 85 85 85 85 85 85	4 48 4 46 6 61 6 61	500 725 640 465 625 2,740 825	\$5555555	20 22 23 19 19 6.1	84 44 58 58 58	8288378	55 58 58 58 58 58	950 3,265 1,525 2,825 965 1,025	850 3,240 1,025 1,325 3,240 690	44 15 4.0	130	шшш	H 150 150	355 450 475	305 440 440
1					6.83	TIELL	ED A	SHIELDED AND REINFORCED	EINE	DRCE										
A   230   Z   Asphalt	-		3 57	23	51	46	570	555	2.5	47	56	7.07	11 700 11 998	1 998 1	101	- 00 -	1	1	-	

Table 74.—Averages of maximum pit depths on coated and corresponding dare pipes with soil resistivities—Continued.

		Cooting		17	IV League City,	ijţ.	Tex.				Pre	V Preble, Ind.	_			υ	ouncil	VI Council Hill, Okla.	kla.	
		- Street		Pit d (m	Pit depths (mils)		Resi	Resistivity		Pit	Pit depths (mils)		Bes	stivity		Pitd	Pit depths		Rosie	Linit
-	ľ		Line	e e	Ş	Control	(ohr	n-cm)		Line	ి	Control	ф	(ohm-em)	Li	Line	Co	Control	uqo)	(ohm-cm)
ssendpidT (slim)	Associated griteos	Character	Coated	Иевтевт Бате	opisogqO gaitaoo	Opposite parest orad	erisoqqO gairoo	etisoqqO Jeorgen orad	Conted	Иеатенt Баге	opisoqqO gailago	erisoqqO Jestaen erad	elieoqqO gaitaoe	otisoqqO tsorsan orad	Coated	Nearest oard	olisoqqO gailaoo	optisoqqO testast orad	orisoqqO unirsoo	etisoqqO tsensen end
		1	COI	D AP	PLIC	ATION	S: CL	COLD APPLICATIONS: CUTBACKS,	KS, C;	AND :		EMULSIONS.	V.S. A							
65	<b>→</b>	Cutback coal tarAsphalt emulsion	42	33	76 71	69	1,400	1,750	46	35	50	37	2,200	3,000	38 38	79	59	##	3,200	3,100
							2. E	ENAMELS	STS											
8888 :	 □ : □ :	Coal-tar-asphalt enameldo do Coal-tar enameldo							113	35	62	8 : :	60 2,900	3,000	8888	488 488 488	96 84 14 14 14 14 14 14 14 14 14 14 14 14 14 1	55 ± 45 52 52 52 52 52 52 52 52 52 52 52 52 52	4,100 3,150 2,450 1,650	5,100 3,250 2,600 4,750
							3.	3. MASTIC	ري											2
519		Asphalt mastic	0.2	51	90	95	1,350	1,350   1,900  0.05	10.05	35	49	9	15,500	15,500 [3,000						
- 1		4. SHIE	LDEI	CO	TING	SHIELDED COATINGS: CUTBACKS,	TBAC	KS, C;	; EMI	EMULSIONS,		A; AND ENAMELS	ENA	TELS						
863 819 819	HZPC	Cutback asphalt. Asphalt emulsion. Coal-tar-asphalt enamel. Coal-tar enamel.	19	52	84	84 108	1,650	1,650   1,500 1,400   1,750	7.4	34 34	50	844	2,250 3,500 2,500	3,900 2,250 2,250	43 11 11 11 11	4444	27,00	8888	3,900 3,750 5,000 2,600	2,350 5,500 4.100
						5. RE	INFOR	REINFORCED	COATINGS	INGS										
107 150 151 143 171 171	M M	Grease Asphalt do do Asphalt enamel Coal-tar-asphalt enamel Coal-tar enamel	-3k-33c3	37 37 37 37 37	11 22 32 32 12 12 12 12 13	15 12 13 13 14 14 15 16 16 16 16 16 16 16 16 16 16 16 16 16	1,750 1,100 1,650 900 1,350 1,000	55 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 8 5 5 5 8 5 5 5 5 8 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5	14 12 8.6 6.5 9.9	39 955	25 4 4 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6	38 39 39 39	5,300 1,950 2,750 4,250 2,900	3,900 3,000 4,000 2,800	25	47	57	4 62 62	2,175	2,450
					6.83	HIELL	ED A	SHIELDED AND REINFORCET	EINF	RCEI								3	2011	20010
230	- Z	Asphalt	2.1	53	65	58	3,000	3.000   1.350   3.8	80	4.5	46	36	1 800	1 800 14 000		-	-		1	

c	Dites	Spindle Top Gully, Tex	Pit depths Daries Pit depths Processing	trol (ohm-cm)	betacc teatrace	Bad Opportunity of the Co.	#		455	1	6   89   H     370   345   0.0   94   89   72   1.090   02=		1.89 79 69 77 30 78 78 78 78 85 88 88 88 88 88 88 88 88 88 88 88 88	Section   Sect
	VII	Caney, Kans,	(mil.	Lane Control (ount-cm)	Nearest bare Opposite coating Opposite Dare Dare Dare Dare Coating Dare Coating Coating Coating Coating Dare Dare bare bare bare	B	2/ 40 54 52 1,700 1,700	2. E	30 68 83 1,400 1,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13,500 13	3. MASTIC	-l 🗅	3 40 69 759 1 2000	5. REINFORCED C	
		Coating ²		- pə	lodmyd Amboldm (alim) Saosiata Sosting O	انا	65   F   Asphalt emulsion	60   G   Coal-tar-asphalt enemal	do Coal-tar enamel	0   519     Asphalt mastic	4,	: : :	107   Canada	150   Checker   150   Checker   150   40   52   44   1.550   1.400     2

Table 74.—Averages of maximum pit depths on coated and corresponding pare

Pit depths														Sites								
Prideptile   Pri							Mt. A	X uburn,	Ħ				Skia	XI took, Ol	da.				Bull	XIII		
Time   Control   Control				C ON WILES		Pit (	lepths iils)		Resi	stivity		Pit (1	depth	on.	Re.	istivity		Pit	depth mils)	es e		stivity
Contacted   Cont	- 1				Ē	ine.	C)	ntrol	편 ()	n-em)	П	ine	ŭ	ontrol	<u>o</u>	ım-em)		Line	) 	ontrol	(o)	n-cm)
28   X   Cont-tar-asphalt enamel   28   35   250   1500	Бутарот	asənəsidT (slim)	Associated gnitsoo		Coated	Nesrest ered	etisoggO gaiteos	etisoqqO teetaen etad		Opposite		Mearest bare	etisoqqO anitaco	ejisoqqO jesnesn	Opposite	etisoqqO jsersen		Teares V.	ətisoqqO	etisoqqO tsətaən	Opposite	nearest
State   Applicate enumerical condition   23   35   57   58   2.256   1.200   1.300				ਜ	8	LD A	PLIC	ATIO	IS: CU	TBACI	ES, C;	AND	EMI	ULSIO								
Con-lar-asphalt enamel   16   66   69   2,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,050   1,05	0-4	21 65	M	Cutback coal tar Asphalt emulsion	23 48	35	70 20		2,250 2,600						1 :::	: :			_		440	_
60 G G Coal-tar-asphalt enamel   23 55 66 68 69 10.00   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.800   1.80										NAME	I.S								-	-		-
State   Asphalt mastic   Asphalt masti	ZMZH	8888	E GO	Coal-tar-asphalt enamel do Coal-tar enamel do	33 44 23	25 25 25	75 68 60 65	69 69 50	1,800 2,050 3,000 1,200			107 87 87	 12 8 12 4	4334 4	265 265 590 590			74	<u>: : :</u>	43	425	<u> </u>
519   State   Asphalt mastic   4. SHIELDED COATHOS: CUTBACKS, C; EMULSIONS, A; AND ENAMELS   4. SHIELDED COATHOS: CUTBACKS, C; EMULSIONS, A; AND ENAMELS   4. SHIELDED COATHOS: CUTBACKS, C; EMULSIONS, A; AND ENAMELS   4. STATE   4									က		2											
Contract saplate enamel		519		Asphalt mastic.							2.8	_	52	_	1,000	_	-	_	_	_	440	465
22   Cutback sapinit curvained   24   66   65   1,600   1,550   1,200   1,550   1,200   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,500   1,50	-			4. SHIE	LDE	8 0	ATIN	38: Ct	TEAC	KS, C;	EMU	LSIO	AS, A	, AND	ENA	MELS						
107   Grease   17   46   65   2,300   1,450   37   102   88   90   365   210   17   92   54   41   380   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   1	HE0H	419 63 81 81 81	LXPC	Cutback asphalt. Asphalt emulsion Coal-tar-asphalt enamel. Coal-tar enamel.	5.1224	85 85 85 85 85 85 85 85 85 85 85 85 85 8	24 68 54	•	2,000 1,600 1,550 1,200								. 6.6 9.0 116		148 141 141 141	4444	375 400 450	
107   Greese   17   46   65   55   2,300   1,450   87   102   88   90   365   2,10   127   92   54   41   390   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150   150								5. RE	INFOR	CED (	COATI	INGS										-1
6. SHIELDED AND REINFORCED   280   Z   Asphalt.	HARREDL	107 150 151 151 143 171 351	:::×:×	lt enamel	17 19	· · · · ·						102 107 107 71 71 71 71	255 559 559 559	90 101 55 48 46 48	<u> </u>	:		: : :	42 41 41 63	44% 4	390 415 440 390	465 390 450 375
230   Z   Asphalt								SHIEL	DED.	AND E	LEINF	ORCE	C)									
	×	230	i			1 1					8 6	26	_	_		_	1 2.6	1	-		006	1

TABLE 14. - Averages of maximum pit depths on coated and corresponding bare pipes with soil resistivities—Conlinued.

	4						ž	Siten	The same of the sa		
					Chamber	XIV Сватветевитя, Ра.			XVI Cholame Flats, Ce	Calif.	
			('outing2	E	Pit depths (mils)		. Resistivity	Pit.	Pit depths (mils)	Resistivity	
				Line	رة -	Control	/mo-ma	Line	Control	(comm-cm)	
Symbol	Thick-	Associ- ated	Character	Control Nearest	est Opposite conting	Opposite nearest bare	Opposite Opposite coaling bare	Coated Nearest	Opposite Opposite	ite Opposite Opposite	est re
:			1, COI.	D APPLICATI	ONS: CUT	BACKS, C	1. COLD APPLICATIONS: CUTBACKS, C; AND EMULSIONS,	ONS, A			
U	228	72	Cuthack coal tar	26 31	59	- 20					1 : :
			To a control to the control of the c		2. EN	2. ENAMELS					
LEHN	88% S	CD H	Coal-tar-asphalt enameldo	2.5 36 21 36	56 84	.00			÷ 13	140	
					3. MA	MASTIC					
0	619		Asphalt mastic					0.39	L	315	1:
			4. SHIELDED	COATINGS:	CUTBACKS	3, C; EMI	SHIELDED COATINGS: CUTBACKS, C; EMULSIONS, A; AND ENAMELS	D ENAMELS			
HCH	819 819 819 819	-LINAC	Cutback asplialt. Aspliat emulsion. Coal-tar-asphalt enamel. Coal-tar enamel.	2.9 32 3.9 28 2.9 33	54 48 	42 51 53		40 81	ТН	140	1::::
		_		Н	REINFORCED COATINGS	ED COAT	INGS				
EXH2	201102	×	Grease. Asphalt. do.		41 48 48	51 53 53		51	153 H	210	: : :
ಜ೯೯	143 171 351	K	Asphalt enamel. Coal-tar-asphalt enamel. Coal-tar enamel.	1.8 35 0.08 32	55	46		20	HH.	<u> </u>	: : : :
				6. SH	SHIELDED AND REINFORCED	ND REIN	IFORCED				1
×	230	Z	Asphalt	2.4   32	48	49		33	П Н	235	1:
1 Ave 2 See 3 Seo 4 Orig	rages ar tables 1- Appendi inal surf	e usually 4 and 15 ix 2 for de face gone,	Averages are usually based on 20 one-foot sections of coated pipe; 14 one-foot sections bare lines; and the equivalent of 2 three-foot sections of 3-in, bare control pipe. See Appendix 2 for description of coatings. See Appendix 2 for description of soils. Coefiginal surface gove, making pit-depth measurements inapplicable.	oated pipe; 14 on napplicable.	ne-foot secti	ions bare l	ines; and the equiv	ralent of 2 three-fo	ot sections of 3-in.	, bare control pipe	å

It was impossible to expose all types of coatings to all the soils, and therefore the coatings were so distributed as to test them in the soils to which they appeared best suited, that is, the heavier coatings were placed in what was thought to be the more destructive soils. For this reason, the summary of the condition of the line pipe under the coatings, table 75, tends to favor the lighter coatings. Strictly speaking, the data in table 75 are not comparable, as all coatings were not subjected to the same soil conditions. As table 74, which averages the pit depths beneath each foot of coating, tends to minimize the seriousness of the corrosion associated with the coating, table 76 has been prepared to show the deepest pit associated with 20 feet of the coatings. Tables 74 and 76 together illustrate the pit-depth-area relation and the extremes of corrosion that may occur under nominally the same condition. Table 76 also shows how difficult it is to produce even 20 feet of coating that will prevent corrosion for 10 years, even when soil conditions are as mild as they were at most of the API test sites. The mildness of the soils is shown in table 74 by the shallow pit depths on the unprotected pipe.

Table 75.—Summary of conditions of line pipe undercoatings.

Metal attack Pitted		deepest pit (mils)		n 322+	+ 822+	264 195 231	* 322+			89		200 107 1 322 +	240		89 290 197 160 197 126 58	124
	ted	Percent		10.00	0.00	62.0 55.0 98.0	70.1		0 6	0.4		70.0 35.9 27.8	48 0		66.3 50.0 50.0 50.0 39.9 37.7	52.2
	Ē	Total feet		139		94 101 148	110		9	9		115 61 56	90		138 125 96 100 105 87 87	,853
	attack	Percent		9.6		10.7 29.6 2.0	4.0		6.6			18.3 21.8 25.3			23.5 23.5 26.5 20.2 23.5 33.1 61.1	22.9
	Metal	Total feet		52 80		975 8 C	3		14		-	30 37 51 9	-	-	61 39 70 105 42 51 58 140	
	Kusted	Percent		5.4		8.8 9.3 8.8 8.8			18.4		-	10.4 39.0 17.9 4.0	-	-	20.00 0.00 114 0.00 0.00 0.00 114 0.00 0.00	0.0
	Kur	Total feet			21 155 0 0			39	ATINGS	-	17 66 36 7	ATINGS	Contract	252 253 253 253 253 253	-	
Inaffortal	מו מעוד	Percent	COLD APPLICATIONS  1.2 0 2. ENAMELS  7.2 15 7.2 15 19.8 6		MAGINTO		72.5	SHIELDED COATINGS		3.5 29.3 43.0	REINFORCED COATTNES	-	0 0 4.7 20.2 30.8 14.9 14.9			
Beall		Total feet	1. (?0]	0.0		21 13 0 31		-	194	4. SHIT	•	59 76	5. REINFO	-	42 42 67 26 12 521	-
	Total feet	inspected		166		152 183 151 157		913		$\int$	164	170 202 177		208	166 192 228 208 218 175 229 3,537	-
Coating	The same a females of the same	(Tharmeter	Cuthnoli	Asphalt emulsion	7	Coal-da enamel Coal-da enamel do		Asphalt mastic			Sphalt amulata	Coal-tar-asphalt enamel.				
	rk- AHBO-	la) coat- ing	\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	<u> </u>	2	н	-			ĺ		Z1	-	5.5	X	ě.
	Sym- Thick-	fol ness (mils)	C 21	-	_	LEW 808 808	-	0 519			_	H & 81			Z 201 R 151 T 171 X 230 Total	A Through pipe.

Table 76.—Single deepest pit on 20 feet of coated line pipe after 10 years of exposure.

[In mils]

R = Ruste M = Metal + = Hole.	a. attac	k.												
Coating							Test	sites a						
symbol	I	II	III	IV	v	V.I	VII	VIII	IX	x	XI	XIII	XIV	XVI
				1. 0	OLD	APPL	ICAT	ions						
C	20 91	 86	322+ 322+	67 87	137 107	39 37	47			51 105		242 128	 54	
					2	ENA	MEL	s						
M K N L.	43 	69 60 45 52	231 195 264 322+		89 	76 52 89 97	57 84 65 60	170 148	R R	60 70 61 76	148 36 58 151	166	45 16 	63
						3. MA	STIC							
o	R	М	U	U	U		R	38	U		R	U		R
				4.	SHIE	LDEI	COA	TINC	3s					
Y F G	40 M 	R	118 211	73  30	M 90 20		R	126 107 45 220	80 M 10 R	45 68 35 35		120 38 46 240	M M 53	103 322+
				5. F	REINF	ORCE	D CC	ATIN	īGS					
B. SE. Z. R. UT	12 100 12 22 51 61 16	38 55 19 50 51 39	183 61 40	39 77 10 36 59 40 58	52 40 45 21 59	50 40  40	37 61  18 67 123 27	290 144 197 120 32	36  30 U U U 32	89  26  38	54 117 69 56 126 27	202 197 132 	23 30 21 23 R	103 80 70 17
			6. SI	HELE	ED R	EINF	ORCE	D CO	ATIN	īGS				
x	1.5	17	22	м	2.4			121	м		31	M	11	124

[&]quot; See table 19 for location of test sites.

U = Unaffected.
R = Rusted.

⁽³⁾ Coatings Applied to Isolated Short Sections of Small-Diameter Pipe.—The results of the tests shown in tables 77 and 78 furnish data on 46 coatings applied to 2-foot lengths of pipe having an outside diameter of 3 inches and supplement Ewing's tests on similar and occasionally identical coatings or coating materials. From a technical point of view, the data on the short lengths of pipe are of interest because they permit a comparison of two methods of testing coatings.

3
nine sections.
)C SC
short
pit on
deepest
) (Jic
1820
Depth of
PABLE 77.
LABI.

	TVI. Cholame Flats, Calif.		83335KK	ç	22 25 27 27 27 27 27 27 27 27 27 27 27	57 70 U
	XIV. Chambers- burg, Pa.	-	RESERE	1	MNDD&MDDMDD	BDDW
	XIII. Bunkie, La.		거=恕====		SCHAHZMADH HORMAZHANDH	2H 109 U
1	XII. Mendota, Calif.		#DD %D # 85 %		#5088480000	0000
i.	XI. Skiatook, Okla.	:	232ZZZZZZZ		RCCCCSSREE	ZZZZ
	X, Mt. Aubum,	-	ZHZZHZHZ		BOGOOMACKEE	CKKR
	IX. Long Beach,		22 23 24 25 25 27 27 27 27 27 27 27 27 27 27 27 27 27		2A22A2454DDD	48××
Test sites 1	VIII. Spindle Top Gully, Tex.		8483528HI		2552 M 4 4 4 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	GZZD
•	VII. Caney., Kans.		24 32 32 32 32 32 32 32 32 32 32 32 32 32		USB 38 3B CB C	<b>314</b> 5
	VI. Council Hill, Okla.	22	222222222		R 223 314 16 27 20 20 20 20 20 20 20 20 20 20 20 20 20	%≅>>
Ÿ	V. Preble, Ind.	ATTON	MMZ1228	వర్త	AGDASS SECOND	UCM 29
	IV. League City, Tex.	APPLICATIONS	M822 22 22 22 22 22 22 22 22 22 22 22 22	ENAMELS	22 R 24 11 14 14 14 14 14 14 14 14 14 14 14 14	2000
	III. Beaumont.	COLD A	101 101 101 101 101 101 101 101 101 101	2. E		ZSH.
	II. Arkansas City, Kans.	1. C	222222 222222		M80008M2 :UMU M	ZHZ
	I. Temple,		REENTE E		HODHATARRAM &	\$4DD
Conting *	Character		Cutback asphalt do do do do do do Asphale emulsion do		Asphalt Coal-lar asphalt Coal-lar asphalt Coal lar do Asphalt	do. do. do.
-	Анко- сіяце совыт 4	-	o hib ppp f, ff		www ww ss kk d, ddd p, zz, zzz zz, zzz zz, zzz, g	
	Thick- ness (mils)	Ī	212 202 213 213 215 215 215	-		337
1	Nym- bol		ec u l b b b ppp fff		VV W W W W W W W W W W W W W W W W W W	uu uu A

	HEDDD		M22M28888 4 7223M2D		HK28K
	GEGEG		%4AAA8AAAbAbb		MIDE
	Zűbbb		K28KH255H8880		17 13 33
	ಜಿಷರಿದಿದ		<b>X&amp;XD#3&amp;XDXDXD</b>		Score
	AMCDD		MM2262MM20MMDD		550 54
	#MM#p		dere Hereit		SMAR S
	MDDD		KUZZEZZZEZZDZ		MM 100+
	880EM		N882193831218833		20 25 15 37
	MADDD		ZHZZZZZZZZZ	gs	ZZZZ
æ	MAMba	gs	REINSHSMENNAR	6. SHIELDED, REINFORCED COATINGS	MM2
SHIELDED COATINGS	Medde	5. REINFORCED COATINGS	PRZZEZZZZZ	CED C	RMMS
co co	MMDMD	ED C	KKKKSKHKKKHKSS	NFOR	41 14 M
TELDI	122 63 0 U	NFOR	2112 274 274 275 274 275 275 275 275 275 275 275 275 275 275	d, REI	35 38 2H
4. SH	MADPP	5. REI	REMERCALES RATES	ELDE	2K2K
	*****		ZHZZZZZZZZZZZZ	6. SHI	ZZZZ
	Cutback sapitalt. Asphalt emulsion. Coal-tar-saphalt enautel. Coal-tar-enamel.		Cutback coal tar Cutback coal tar Asphal, emulsion Grease Grease Asphal, Asphal, do do do do do do Coal-tar-asphall enamel		Grease Ashlate do Metal coating
and the state of t	f, fiff B d, dd g, p, zzz		cc b ff, fff e e e e e e e e e e e e g o t, g pp, qq y, yy y, yy k k k		5
li dana	18 190 8 41 8 50 8 77		22 22 22 26 26 27 26 26 26 26 26 26 26 26 26 26 26 26 26		105 199 216 0.4
i	r ff ss ddd		d kk		r y g y

In mile.
See the 15 for a description of the coatings.
See Appendix 2 for a description of the soils.
The sessented coating is similar to the soils.
The restricted coating is similar to the coating on the same line except for a change in priming coat, reinforcement or shield.

Rusted—no appreciable pit depths.
 Pits less than 10 mils in depth.
 Pipe shows no corrosion.
 Without shield.

TABLE 78.—Summary of conditions of 3-inch coated pipe.

Total pipe   Percent	Total pipe   Total pipe   Percent   Total pip	Conting			Unaffected	seted	Rusted	ાળ	Metal attack	attack	E	Pittad	
1. (U(1) APPLICATIONS	1. (Vol.) Applications  2. (Vol.) Applications  3. (Vol.) Applications  4. (Vol.) Applications  4. (Vol.) Applications  4. (Vol.) Applications  4. (Vol.) Applications  3. (Vol.) Applications  4. (Vo	<b>5</b>	aracter	Total pipe inspected	Total pipe		Total pipe		Total pipe	Percent	Total pipe	i	Depth deepest pit (mils)
15 0 0 0 4 26.7 6 40.0 5 83.3 8 80.0 15 15 0 0 0 0 1 1 6.7 8 10.0 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 183.3 8 18	15 0 0 0 1 1 26.7 8 40.0 8 53.3 8 1 15 1 15 1 0 0 0 1 1 1 26.7 8 40.0 8 53.3 8 1 15 1 1 1 20.0 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1				1. 00	LD APPLI	CATIONS						
2. ENAMELS  15 0 0 0 2 13.3 6 40.0 7 46.7 115 7 46.7 115 7 46.7 115 7 46.7 115 115 115 115 115 115 115 115 115 11	2. ENAMELS  15 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	Cuthack asphalt. do. Cuthack coal tar do. Asphalt emulsion do.		55555555	00101000	0 6.7 6.7 0 0 0 0	7#110010	26.7 20.00 6.7 6.7 6.7 0 0 0	\$000 #A04	20.0 20.0 40.0 13.3 26.7 6.7 6.7 6.7	28720121	83. 46.7. 86.7. 93.0. 93.0.	2 2 2 2 2 2 2 2 2 2 2 2 2 2 3 2 3 3 3 3
15   0   0   0   13   3   6   6   0   0   14   15   15   16   16   17   17   17   17   17   17	15   0   0   0   0   13.3   6   40.0   7   46.7     15   7   46.7   2   13.3   6   40.0   7   46.7     15   4   26.7   2   13.3   6   40.0   4   20.7     15   4   26.7   2   13.3   6   40.0   3   20.0     15   4   26.7   2   13.3   6   40.0   3   20.0     16   4   26.7   2   13.3   2   14.2   3   2   14.2     15   7   46.7   3   20.0   1   6.7   3   20.7     15   7   46.7   3   20.0   1   6.7   3   20.7     15   3   20.0   1   6.7   2   13.3     16   3   20.0   4   26.7   8   53.3     17   46.7   3   20.0   1   6.7   8   53.3     18   53.3   3   20.0   0   0   0   0     19   66.7   2   13.3   3   20.0   0     10   6.7   2   13.3   3   20.0   0     11   12   13   2   2   2   2   2     13   3   2   2   2   3   3     14   5   5   5   3     15   5   5   5   3     16   7   7   7   7     17   7   7   7     18   7   7   7     19   7   7   7     10   7   7   7     10   7   7   7     11   7   7   7     12   7   7   7     13   7   7   7     14   7   7   7     15   7   7   7     16   7   7     17   7   7     18   7   7     19   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7     10   7   7						LS						:
	3. MASTICS 3. MASTICS 3. MASTICS 15 2 13.3 1 6.7 2 13.3 10 66.7 1 15 10 66.7 1 15 10 66.7 2 13.3 20.0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	Asphaltdo.coal-tar asponder asponder asponder coal tardododododododo	had to	22222224222	007401407700	0 0 0 0 0 0 0 0 0 4 26.7 42.9 46.7 46.7 46.7	3033111200	13.3 0.1 13.3 14.2 20.0 20.0 20.0	C4001400101	0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.0	ь <u>П</u> 42200ги401	28.7.2 20.7.2 20.0.7 20.0.7 20.7 4.0.0 4.0.0	20 H % S H 22 L S H 90 L
	,	Asphaltdododo.		15 15 15	10 83 10	13.3 20.0 53.3 66.7	2301	6.7 0 20.0 13.3	01400	13.3 26.7 20.0 20.0	10 8 1	66.7 53.3 6.7	H H 30

	H 63 212		21H852HHH853H8%H		88 38 H	
	46.7 26.7 6.7 0		0.85.0.0.0.0.0.0.0.0.0.0.0.0.0.0.0.0.0.0		33.3 60.0 13.3 100.0	
	74100		######################################		152295	
	33.3 46.7 13.3 13.3		8100 88 88 88 88 88 88 88 88 88 88 88 88 8		53.33.33 53.33.33	
	DDD0-101		222114228925 2225		ω14ω0	
	13.3 13.3 6.7 6.7		6.7 13.3 13.3 13.3 6.7 7.7 7.7 6.7 0 0 0 0 33.3	TINGS	13.3 0 0.7 0	
OATINGS	20111	5. REINFORCED COATINGS	-40040-4-0-0066	SHIELDED, REINFORCED COATINGS	2010	
4. SHIELDED COATINGS	6.7 73.3 80.0 80.0	TORCED	0 0 0 0 0 0 15.4 13.3 13.3 13.3 20.0 20.0 20.0	, REINFOI	0 6.7 26.7 0	
4, SHI	12 112 123 123	5. REIN	000	HIELDED	0140	
	15 15 15 15		22222222222222	6.8	15 16 15	
,	Cutback asphalt. Asphalt emulsion. Coal-tar-asphalt enamel. Coal-tar enamel. do.		Cutback asphalt Cutback coal far Asphalt emulsion  do do Asphalt do do do do do do do do Coal-iar-asphalt enamel Coal-far enamel		Grease. Asphalt Asphalt Metal coating.	
	f, fff s d, dd zzz, p, g		fr, fff e ve vy, yyy yy, yyy y, yy y, yy k dd, ddd		y, yyy q	hiold
	18 190 141 150 177		72 894 149 149 155 155 165 165 165 342 342		105 190 216 0.4	1 Without shield
	r ff ss ddd zz		bb Pb Pb Pb Pb Pb Pb Pb Pb Pb Pb Pb Pb P	Í	yy yx dag	1 Wi

(4) Comparison of Results of Tests of Coatings Applied to Operatina Lines with Those on Isolated Small Pipe.—Table 79 was prepared for the purpose of showing the relative severity of tests of coatings applied to pipe lines and to short isolated sections of small-diameter pipe. The data are from three sources: Conductance tests, pattern tests, and measurements of pit depths. The data were derived as follows: If the results of each of the four conductance measurements made on the line coating were lower than the result of the single conductance test of the coating on the small pipe, the number of the test site for which the comparison was made was placed under the heading "lower on line." If the conductance of the coating on the small pipe was lower than any of the four coating conductances for the coating on the line, the site number was placed under the heading "higher on line." If the conductance of the coating on the small pipe was less than the conductance found by one test of the line coating and greater than that found by another test of the same coating, the site number was placed under the column headed "indeterminate." The results of the pattern tests were treated similarly. The comparison of pit depths was based on the average of the depth of the deepest pit on each of about 20 one-foot lengths of coated pipe line and the depth of the deepest pit on the small pipe similarly protected. The columns under the heading "pit depths" group the test sites with respect to whether the averages of the maximum pit depths on the line beneath the coatings were less, greater, or equal to the maximum pit depth on the corresponding small pipes.

Table 79.—Comparison of results of tests of coatings applied to pipe lines and isolated sections of small-diameter pipe.

							==
Sym-	Thick-			Conductance test			
bol	ness	Lower on line at sites	Σ	Higher on line at sites	Σ	Indeterminate at sites	Σ
		1.	COI	LD APPLICATIONS			
C A	21 + 65 +	VII.	1	ш, х	2 0	I, IV, V, VI, XIII XIII, XIV, I, II, III, V, VI, X	5 8
				2. ENAMELS			
N K M L	60 + 69 - 58 + 80 -	VI, XI	1 0 2 1	II, VII, VIII, XI II, III, VI, VII, XI, XIII, XIV X. XIV II, VIII, IX, X, XI, XVI	7	III, VI. I. III, VII. III, VI.	2 1 4 2
				3. MASTIC			
Ů	519 -	II, III, IV	3		0	I, V, VII, VIII, IX, XI, XIII, XVI	8
		4.	SHI	ELDED COATINGS			
Y F G H	29+ 419+ 63+ 81+	IX. X. XIII	0 3 2 3	VI, VIII, IX, X, XIII. I, VI III, VIII, XIII, XVI III, IV, VI, VIII, XIII	5 2 4 5	I, IV V, VIII, XIII, XIV, XVI II, V, VI, IX	2 5 4 1

Table 79.—Comparison of results of tests of coatings applied to pipe lines and isolated sections of small-diameter pipe—Continued.

·	Thick-	1		Conductance test		
bol	ness	Lower on line at sites		Higher on line at sites		Interminate at sites
		5. I	REIN	FORCED COATINGS		
B S E Z R U	107+ 150+ 151- 201+ 143- 171+ 351+	VII, XI IV XIV, XVI V, XIV, XVI III, IV, IX, XI, XVI III, IX, XIII, XIV, XVI XIII, VII	2 1 3 5 1 5 2	II, V, X I, XIV VI, X VI, X II, V, VIII II, V, III, IX, XI, XIV IV, VI, X, XI, XVI	3 2 2 4 5 1 5	I, IV, VI, IX, XIV, 5 II, VII, VIII, XIII, V. 5 I, II, XI, XIII, IV 5 VII, XIII 2 IV, V, VII. 3 II, IV, VIII, VIII, XI 5 I, VIII, IX 3
		6. SHIELI	)ED	REINFORCED COAT	rinc	ss
X E	230+	I, IV, V, XIV.		IX, XVI	61	II, VIII, III, XI, XIII 75
G	mhi-l- l			Pattern test		
bol	Thick- ness	Better on line at sites		Worse on line at sites		Indeterminate at sites
		1.	COI	LD APPLICATIONS		
C A	21+ 65+		0		0	I, III, IV, V, VI, VII, X, XIII, V, VI, XI, XIII, XIV, Y,
				2. ENAMELS		
N	60+		0	ır	1	III, VI, VII, VIII, IX, XI
K M L	69 — 58 + 80 —	IX, X	0 0 2	II, III, XI, VI, VII, XIII, XIV	7 0 0	Y I, III, VI, VII, X, S II, III, VI, VII, VIII, 7
				3. MASTIC		
0	519+	II, XIII, IX.	3	IV, VII.	2	I,III,V,VIII,XI,XVI 6
		4.	SHI	ELDED COATINGS	·	
G H	63 + 81 +		0	II, III, IX, XVI, VIII, XIII III, IV, V, VIII, X, XIV	6 6	VI, V, X, VII
-		5. R	EIN	FORCED COATINGS		
B S	107+ 150+		0	X, XI X, XIV	2	I, II, IV, V, VI, VII, S I, II, IV, VII, VIII, S
E Z	151 — 201 +		0	I, II, X, XIV I, IV, V, VII	4 4	II. II. VIII, VIII, KIII, XXIII IV. VIII, XXIII XVIII, XXIII, XXIII
R U	143 — 171 +		0 0	I, II, IV, V, VII, IX, XI, XIV I, IV, VII, XIV	8 4	III, VII, VIII, IX 4 II, III, XI, VIII, XIII,
T	351+		0	IV, IX, X, XI, XVI	5	I, VI, VII, VIII
		6. SHIELI	ED,	REINFORCED COAT	rinc	S
Σ			5		48	93

Table 79.—Comparison of results of tests of coatings applied to pipe lines and isolated sections of small-diameter pipe—Continued.

				Pit depths			
Sym- bol	Thick- ness	Shallower on line at sites	Σ	Deeper on line at sites	Σ	Equal at sites	Σ
Management Surveyor		1.	COI	LD APPLICATIONS			
C A	21 + 65 +	VI, VIIIV, XIII	2 2	I, IV, V, XIII. I, II, V, VI, X, XIV.	4 6	III, X	2 1
	·			2. ENAMELS		·	
N K M L	60 + 69 - 58 + 80 -		1 0 2 0	II, III, VI, VII, VIII, III, III, VII, VIII, XI, XI, XII, XI	7 9 5 9	· iii	0 0 1
		<u></u>		3. MASTIC		<u></u>	
0	519+	III, IX	2	I, II, IV, V, VII, VIII, XI, XVI	8	XIII	1
		4.	shi	ELDED COATINGS		'	
Y F G H	29 ÷ 419 + 63 ÷ 81 +	IX	0 1 0	I, IV, VI, VIII, X, XIII, XIV IX I, V, VI, VIII, X, XIII, XIV, XVI, X, XIII, VIII, IX, X, XIII, VIII, IX, X, XIII, IX, V, V, VI, VIII, IX, X, XIII, XIV	8 8 10 9		0
		5. I	REIN	FORCED COATINGS			_
B S E Z R U		XIII. I, IV, XIII. III. I, III. IV, IX, XIV, III.	1 1 3 1 2 4 0	I, II, IY, V, VI, VII, I, II, IX, X, XI I, II, IX, X, XI II, V, V, VII, VIII, II, XI, X, XI II, IX, XI, XVI II, IX, V, VII, VIII, II, IX, VII, VIII, XI, II, IV, VI, VIII, VIII, II, IV, VI, VIII, VIII, IX, X, XI, XVI  REINFORCED COAT	9 7 7 9 7 6 9	XIII	0 0 1 1 1
<u>x</u>	230+	III, VIII, IX, IV	4	I, II, V, XI, XIII, XIV,		i S	
Σ	<u> </u>	1	26	XVI	8 145		8

The results of the two methods of testing coatings, that is, on operating pipe lines and on isolated sections of small pipe, are not strictly comparable for several reasons.

2. The results of four or more observations are compared with the

^{1.} The thickness of the coating on the line and on the corresponding small pipe was not always identical, as is shown in the second column of table 79, in which a plus sign is placed after the thickness of the coatings that were thicker on the pipe line and a minus sign after the thicknesses of coatings that were thinner on the pipe line.

result of a single observation, the reproducibility of which cannot be determined.

3. The tests were all based on some measurement of a worst condition, and on such a basis, the tests are more severe with respect to the coatings on the line, because these coatings had greater areas from which the worst condition was selected. This statement holds for the conductance tests, although the results were expressed as conductance of 1 square foot of coating, as the conductance was not uniform but was largely controlled by the conductance of some small, defective area. In the case of the pit depth comparisons, the area of 1 linear foot of line pipe ranged between 201 and 338 square inches, whereas the area beneath the coating on the small pipe was about 140 square inches.

Each of the tests, therefore, would have a tendency to make the coating applied to the line appear to be in the worst condition even if the actual conditions on a unit-area coating on the two sizes of pipe were identical.

This is but another way of saying that if the coating is judged by the worst condition found on that coating, the severity of the test increases with the area of the coating exposed to the test. This may be a sufficient explanation of the fact that table 79 indicates that, on the whole, the conditions of the coatings on the pipe line were worse than the conditions of the coatings on the small sections of pipe. This explanation is supported by the large number of cases in which the result of a test of a coating on a small pipe fell between the results of the four tests of the corresponding coating on the operating line.

Table 79 shows that in many cases the coating on the small pipe was in a worse condition than the corresponding coating on the operating line. This condition is not confined to any one soil or coating, and there seems to be no correlation between the relative merits of the coatings in the two tests and either the character of the soil or the nature of the coating. Probably many of the apparent differences in the conditions of the same coating under the two types of test are the results of variations in the conditions of the soil or coating or of other accidental variations. This is suggested by the fact that in both the conductance and pattern tests a large percentage of the results falls in the indeterminate classification.

Although table 79 indicates that, in the majority of cases, the coatings applied to the pipe line deteriorated more than the corresponding coatings applied to small-diameter pipe, the fact that the conditions of the two tests were not identical prevents the conclusion that table 79 shows that a test of the coatings on the line is necessarily more severe than the other test if the exposed areas of the coatings are the same.

A question of equal importance is whether or not the two tests place coatings in the same relative positions. Inasmuch as the plans for the tests specified that the names of the makers of the line coatings should not be revealed, the publication of a table comparing the results given in tables 75 and 78 is not permissible. However, such a table has been prepared for the purpose of answering the question just mentioned. It will be seen that tables 75 and 78 permit the arrangement of the coatings in order of merit on three bases: (1) percentage of unaffected pipe, (2) percentage of coatings that permitted the formation of pits, and (3) the depth of the deepest pits beneath each coating.

When coatings were arranged in order of merit as indicated by the percentage of protected pipe unaffected, percentage pitted, and the

maximum pit depth for the two kinds of exposures, it was found that the three bases for comparison did not arrange the coatings in exactly the same order. The average range in rank for the line coatings in the three tests was approximately 5.4 positions, whereas the average range for the same coatings on the small pipe was about 3.6 positions. Table 80 shows comparisons of the ranks of coatings on pipe lines and isolated pipes grouped according to their structures. The comparisons are based on three criteria of the condition of the pipe beneath the coating, as shown in tables 75 and 78. Table 80 may indicate that the line tests of the unreinforced coatings, which in this table included cold applications. enamels, and a mastic, were on the whole somewhat more severe than the tests of these coatings on the isolated pipe, but the indications are not very definite. The grand totals of the results show that, although two ways of testing coatings yield different results with respect to ranking individual coatings, neither method greatly favors any class of coatings. It seems probable that the differences in ranking by the two test methods are not greater than the variations in ranking resulting from the lack of control of the test conditions.

Table 80.—Comparison of the ranking of coatings on pipe lines and isolated pipe.

	isola	er of linted coa	tings	Number ran	of line ked hig (better)		Number rar	of line aked lov (worse)	wer
Test criterion	Unrein- forced	Rein- forced	Shielded	Unrein- forced	Rein- forced	Shielded	Unrein- forced	Rein- forced	Shielded
Percent unaffected Percent pitted	1 3 1	5 1 3	1 1	1 1 2	2 4 2	1 3 3	5 3 4	0 2 2	3 1 2
TotalGrand total	5	9	2 16	4	8	7 19	12	4	6 22

As the tests were conducted, the conditions were apparently more severe for the coatings on the line, and the results of the tests of coatings on the pipe lines were more dependable because of the greater number of observations for each coating. But these facts do not indicate that tests of coatings by applying them to working lines are better than tests of coatings on isolated sections of pipe for determining the relative merits of coatings if the same amount of data is obtained by each test.

(5) Effect of Time on Rate of Corrosion of Coated Pipe.—It should be evident from the data presented that the production of even a few feet of coating that would remain continuous for as long as 10 years was a difficult undertaking. Data on more recent coating installations indicate that the problem is still difficult. Nevertheless, complete protection of the pipe is the goal sought. A combination of a protective coating with cathodic protection might be more economical and more effective than an attempt to maintain a continuous coating. The current required for cathodic protection will be determined to a considerable extent by the conductivity of the coating. For this reason, rather than as a means of indicating the effectiveness of the coatings, table 81 is reproduced from the final report on the API tests [61].

It should be pointed out that the conductivity of the coating as reported depends largely on the conductance of the solution in the pores and other openings, which in turn depends on the number and size of the openings, the thickness of the coating, and the conductivity of the soil solution in the openings. The data are also affected to some extent by the way in which the measurements were made. Those readers who are accustomed to think in terms of resistance rather than conductance can reduce the conductances to resistances by dividing the latter into 10⁶. It will be seen that some of the coatings are much poorer conductors than others.

It has been shown that in aerated soils the rate of penetration of bare iron and steel pipe decreases as the period of exposure increases. It is of interest to inquire whether the penetration of coated pipe is similar. The API tests do not answer the question positively but throw some light on it.

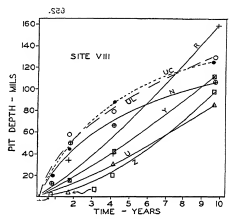


FIGURE 40.—Relation of pit depths on coated pipe to duration of exposure.

UC, unprotected control pipe; UL, unprotected line pipe; R, asbestos-felt-reinforced asphalt enamel; N, coal-tar enamel; U, asbestos-felt-reinforced coal-tar enamel; Y, aluminum-foll-shielded asphalt cut-back; Z, asphalt reinforced by two layers of asphalt-saturated cotton fabric. See appendix 2 for description of the test sites.

Because of the effectiveness of some of the coatings, the mild corrosiveness of the soil of some of the test sites, and the lack of uniformity in some of the soils and coatings, many of the sets of observations did not yield data that definitely indicated the relation of pit depths to time. However, 9 of the API test sites yielded fairly satisfactory data for 3 or more coatings, and these data have been plotted in 12 sets of curves [77], one of which is shown in figure 40. In these figures, pit-depth—time curves were shown for pipe protected by 17 types of bituminous coatings, which were representative of most of the coatings in use when the API tests were started in 1930. Each point on the curves for protected pipe represents the average of the depths of the deepest pit on each of two 5-foot sections of coated pipe in an operating pipe line, usually a line of 8 inches in diameter. The points on the curve for the uncoated pipe line, UL, figure 40, are based on averages of the deepest pit on each of twelve 4-foot sections of uncoated pipe line. The points on the curve for the control pipe, UC, are based on the averages of the

Taber 81,—Maximum conductance of coatings on operating pipe lines.

	To be seen a service of the service	VI. Cholame		08:		변호 년 	777,000	330
		IV. Chambers-	X	E 570				
	1	III. Bunkie, La.	z	E E E E E E E E E E E E E E E E E E E		問因		8 g
		II. Skiatook, Okla.				4,700 1,400 3,500 1,000 17,000 4,700 3,400		32 290
		Z. Mt. Aubum,	:	22 22 23 24 24 25 25 26 26 26 26 26 26 26 26 26 26 26 26 26		D E E E E E E E E E E E E E E E E E E E		
unes.		IX. Long Beach,				23 98 98 8,700 8,700		0.23
g Pupr	E	VIII. Spindle Top Gully, Tex.	1 -4			전혀 : : : : : : : : : : : : : : : : : : :		55 620
he reserve	Test sites	VII. Caney, Kans.	ULSION	4,800		240 1,600 120 E E E E E E 1,100		8 8
of consents on operating pape times.		VI. Council Hill, Okla.	C; EM	4,300 930 E		272 270 270 28 200 30 30 30 30 30 30 30 30 30 30 30 30 3		
		7. Preble,	FBACK	लललल	ELS	E 67		13
		IV. League City, Tez.	NS: CU	2,700 1,600 4,000 4,800	2. ENAMELS		3. MASTIC	12
		III. Beaumont, Tex.	CATIO	<u>न</u> संसम्	23	138 : EE :	eri .	112
		II. Arkansas City, Kans.	1. COLD APPLICATIONS: CUTBACK C; EMULSION	전면		2.8 2.8 2.8 2.8 2.8 2.8 2.8 2.8 2.8 2.8		10
· )		I. Temple, Tex.	1. COLI	2,510 E E E		, DEE:		5 4 110
		See- tion		ಇರಿ ಇರ	Ì	ಇರ ಇರ ಇರ ಇರ		etQ
The second secon	Coating 2	Character		Cutback coal tar		Coal-tar-asphalt enamel do do do		Asphalt mastic
	රී	Asso- ciated coat- ing		Þ ₽	-	о		¥ ::
-		Thick- ness (mils)		21 65	ŀ	69 58 80 80	-	519
		Sym- bol		D ◀		LKKN	-	0

4. SHIELDED COATINGS: CUTBACK Y; EMULSION F: AND ENAMELS	
IELDED COATINGS: CUTBACK Y; EMULSION F: A	ENAMELS
TELDED COATINGS: CUTBACK Y; E	AND
TELDED COATINGS: CUTBACK Y; E	<u> </u>
IELDED COATIN	EMULSION
IELDED COATIN	×
IELDED COATIN	CUTBACK
TELDE	ž
TELDE	ဗ
	TELDE

			<b>-</b> 53		
			1,600 E		:
	2	1,200 1,200	750	13	0.00
	冠	13,300			E
			: :	0.016	
	E	1,700 E	8 9	0.016	4.6
	图	310 810	98	0.87	F. 9
	E	হালে ই	061	S ES ES	±1
,	:		0.005	0 15 E 0 004	: :
	E.	3,600,5	5 62 5	E E	000
	006'8	1,200	0.08	0 15	700.0
	8,900			800 1,800	20042
	-		 000 000 000 000 000 000 000 000 00		
	i	: :	2.5	3 :	-
	ল্ল	220	· !		
	8.4	: a.c	2.5		_
	Cutback asphalt	Asphalt emulsion	Coal-tar-asphalt enamel	Coal-tar enamel	
	బ	Ą	Z	П	
	50	419	63	81	
-	<b>&gt;</b>	r ₌ ,	7 M	н	The same of the sa

COATINGS
REINFORCED
r.

			00	£8	00	530	2,000 2,000	8
	E3	-: -: 28						
			:			O.22E	0	:
		2	17,000	2,900	<b>E</b>	360	800 300 300	
	Q	2,700	6.800		2,20 300 300		00 X 5	0.4
	田	¥1 :	(A)	:			110	:
	2,800	000,86	1,700		2,300		725	
		E	1,700	E	351	2,500	310	
	98	386	4	97	325	220	2 2 2 2 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3	
	터	3 :	E	4			150	
	田	12.8	31212			8		
	터브	i D Š	38	22.2	1,300	1 17	188	
	-			800	98,5	385	3	
	670	230	55	E	2,000	285	3	
ľ	田田	医氏	800	220	땅으	200	8.5	
-	a,0	etC	-8-2	- a.c	8.4	a,c	4.0	
	Grease	alt	do	do	Asphalt enamel	Coal-tar-asphalt ename	Coal-tar enamel	
-	. Grea	. Asphal	<u>:</u>	:	. Asph		. Coal.	1
L			<u>:</u>	×	<u>:</u>	M		
	101	150	151	201	143	171	351	
-	В	ĽΩ	闰	Z	æ	D	H	

6. SHIELDED, REINFORCED ASPHALT

	-
530 4,000	
680	erial,
:	table 16. contain, with the exception of coatings $G$ and $X$ , the same bituminous material,
530 10,000 1,000 5,200	e bitumi
530 14,000	the sam
	and Y,
	oatings (
20	tion of e
43	о ехсер
E 280	ı, with t
40	3 contair
3 6	column
	able 16, ating in
Asphalt	In micro-mloss per square foot, A description of these coatings is given in table 16. The conding in column 1 and the related coating in E pipe exposed. Coating damaged by tools.
Z	hos per on of the g in col posed. maged
230	n micro-mho description The coating i = pipe expo
×	Thur TA de The The Coal

deepest pit on 24 individual 18-inch lengths of pipe. In addition to the tests of coatings applied to pipe lines, the API tests included 46 varieties of coating applied to 2-foot lengths of boiler tube. These specimens were placed in a trench near the pipe line at each site. Only three specimens of each material were provided for each test site. The heavily coated specimens showed no pits at the first inspection. The lightly coated specimens in the corrosive soils were punctured at the close of the test. On this account, these specimens yielded few data from which pit-depth-time curves could be plotted.

Figure 41 shows the data for one test site. The curves are similar to those for the coated line pipe. Incidentally, it is of interest to compare the curves for the uncoated boiler tube, UB, buried with the coatings with that of the uncoated control pipe, UC, buried next to the pipe line

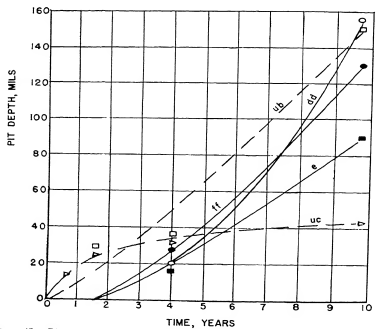


FIGURE 41.—Pit-depth-time curves for coatings applied to boiler tube at American Petroleum Institute site, XIII.—Miller clay.

dd, Coal-tar enamel; e, cotton fabric reinforced, chromated grease; f, asphalt-chromate emulsion; ub, uncoated boiler tube; uc, uncoated control pipe in adjacent trench with pipe line.

a few feet distant. During the first 4 years the two pitted at approximately the same rate but at the end of the 10 years of exposure, three of the four boiler tubes were punctured, whereas the control pipes showed little increase in pit depths. At this site the pitting on the line was much deeper than on the control pipes.

Most of the pit-depth-time curves for the coated pipes do not bend toward the abscissa as much as do the corresponding curves for the unprotected pipe. In a number of cases, the curves indicate that the rate of pitting did not decrease as the exposures lengthened and in several cases there appears to have been a slight acceleration of the rate of pitting. If these curves truly represent the course of corrosion of coated pipe, in the long run many of the pipes that were coated would be punctured before the adjacent uncoated pipe. There are a considerable number of cases in which at the 10-year period of inspection the pits on the coated pipe were deeper than those on the uncoated pipe, and at two sites the pipe beneath one or more of the coatings was punctured.

The general dissimilarity between pit-depth-time curves for bare and coated pipe exposed to the same soil conditions cannot be accounted for by characteristics of the coatings or of the soils to which the coatings were exposed, because widely different coatings yielded similar curves, and the curves are similar for widely different soils. The curves for a pipe protected by a specific coating may or may not be similar in different soils, and the factor that determines the shape of the pit-depth-

time curve is not apparent from the available data.

The results that have been presented may represent the normal behavior of pipe protected by coatings or abnormal conditions arising from the way in which the tests were conducted or the way in which the data were treated. The alternation of coated and bare sections of pipe may have set up conditions of differential aeration that intensified the corrosion beneath the coatings and these conditions may have been aggravated by the exposure of additional pipe and the increased aeration of the soil at each period of inspection. In considering this suggestion, one should remember that it is the shape of the curves, that is, the change in curvature with time, and not the magnitude of the pit depths that is the topic to be considered. Comparisons of pit depths on the uncoated line pipe with those on the uncoated control pipe do not definitely indicate that the unprotected line pipe received cathodic protection at the expense of the coated pipe because of differential aeration.

It is possible, however, that the effect of the bare sections on the coated sections of pipe is obscured by other phenomena and cannot be shown by a comparison of the bare sections of the line with the bare control pipe because of differences in area, soil pressure, curvature of pipe, or other conditions. Unless the bare sections of pipe tended to maintain the rate of penetration of the coated pipe and to make the pits deeper, the effect of the bare sections is of little importance with respect to the curves. It is much easier to show that because of differential aeration the bare sections may have caused an increase in the depths of the pits on the coated sections than to explain how the bare sections caused the rate of penetration of the coated sections to be maintained. As the coatings became older, the effect of differential aeration probably decreased because of the increased number of holes in the coating and the greater compactness of the soil around the unprotected pipe. The results of the tests of the coatings applied to isolated sections of pipe parallel those for the coatings applied to the pipe lines and lead to the same conclusions as to the relation between pit depths and the age of the coatings. This indicates that the presence of the bare sections of the pipe did not exert a controlling effect on the pit-depth-time curves for the curves for the coated line pipe.

The difference in the shapes of the pit-depth-time curves for coated and uncoated pipe could be explained in part at least by the assumption that the pits beneath the coating did not start until some time after the tests began. Correction of the ages of the pits for this lag in time of start would yield curves showing higher initial rates of penetration and, consequently, greater bending toward the time axis. There is no way of determining when the pits started, and therefore it is impossible to

evaluate the importance of this explanation.

The two explanations that have been outlined above attribute the shapes of the pit-depth-time curves for coated pipes to conditions associated with the test rather than with the characteristics of the coatings tested. A second class of explanation implies that the results of the tests reveal tendencies which are to be expected whenever a coated pipe corrodes under certain normal conditions. In the case of coatings containing organic materials that decay, conditions may be more favorable for the development of certain bacteria to which corrosion under certain environmental conditions has recently been attributed [78, 79]. A pipe line frequently picks up and discharges currents commonly designated as long line currents, and an imperfect coating may increase the current density of these currents at points where the coating has failed.

The theory of differential aeration as a cause of corrosion, which was mentioned in the discussion of the effect of the unprotected sections of the line, may also be used to explain why sustained rates of corrosion of coated pipes are to be expected. Rates of underground corrosion depend on two factors: the tendency of the metal to go into solution, indicated by the potential of the corroding metal with respect to the electrolyte, and the corrosion-repressing power of the corrosion products including those frequently termed polarization films. Both of these factors are influenced by the amount and distribution of the oxygen in the soil. It has been shown [56] that the poorer the aeration of the soil the more nearly constant is the rate of corrosion. It is reasonable to suppose that the coating reduces the accessibility of oxygen to the pipe and hence tends to maintain a more nearly uniform rate of pitting than on unprotected pipe. This seems to the writer to be the most reasonable explanation of the sustained rates of corrosion shown by many of the pit-depthtime curves.

Since the pit-depth-time curves in figures 41 and 42 represent only selected data and are free-hand curves arbitrarily drawn, it is desirable to eliminate the personal equation involved and to examine all of the pit-depth data for the longest period of exposure to see whether they point to a similar conclusion. Several basically different comparisons of pit-depths on coated and uncoated pipe can be made, although each

comparison has some objectionable feature.

The comparison that will be presented first is based on the data from which the curves were plotted. The final report [61] on the API coating tests shows 30 cases out of a possible 175, or about 17 percent, in which the average of the depth of the deepest pit on each of two 5-foot sections of coated pipe was deeper than the average of the depths of the deepest pit on each of twelve 4-foot sections of unprotected pipe at the same site. The 30 cases comprise 12 coatings and 10 test sites. If the single deepest pit on a section of pipe is made the basis of comparison, there are 67 cases out of a possible 334 cases, or 20 percent, in which the deepest pit on a 5-foot section of the coated pipe was deeper than the deepest pit on the 4-foot section of uncoated pipe nearest to the coated section. The 67 cases comprise 14 coatings and 12 sites.

Maximum pit depths are to some extent the results of additive combinations of circumstances and are therefore less significant than averages, as the latter tend to minimize these cumulative circumstances. There are 16 cases out of a possible 169, or 9 percent, in which the av-

erage of the depths of the deepest pit on each of approximately twenty 1-foot sections of the coated pipe was greater than the average of the depths of the deepest pit on each of about sixteen 1-foot sections of unprotected pipe adjacent to the coated pipe. The 16 cases comprise 7 coatings and 8 soils, and when compared with the data previously given, illustrate the effect of basing comparisons on averages instead of maximum figures.

As the pitting of the bare sections of the line may have been influenced by the alternation of coated and uncoated sections, a comparison will next be made of the pit depths on the coated line with those on the adjacent lengths of isolated control pipe. There were 10 cases out of a possible 156, or 6 percent, in which the average of the deepest pit on each foot of pipe beneath a coating was greater than the average of the deepest pit on each of two 36-inch lengths of control pipe adjacent to

the coating. The 10 cases involve 7 coatings and 4 soils.

The relative slope of the pit-depth-time curves for coated and uncoated pipe is more important than the relative depths of the pits at the close of the 10-year period of exposure. There are 83 cases in which measurable pits were found beneath coatings on the operating lines at both the third and the fourth inspection. If the maximum pit depth on the 5-foot coated section is divided by the average of the depth of the deepest pit on 4 feet of each uncoated section of pipe at the same site, there are 52 cases out of the possible 83 in which the quotient is greater for the last period of inspection than for the preceding one. In other words, in about 53 percent of the cases the rate of pitting beneath the coatings decreased more slowly than the rate of pitting of the uncoated pipe. This comparison is too favorable to the coatings because it does not take into account 52 sections of coatings that prevented pitting during the first four years of exposure but permitted pits to develop during the subsequent 6 years.

The comparisons of pit depths on coated and uncoated line pipe tend to confirm the indications of the pit-depth-time curves. As the data for the pit-depth comparisons were derived from the same source as those for the pit-depth-time curves, these comparisons show only that the curves have been plotted correctly. Because the data presented above may have been affected by the interaction of the coated and uncoated sections of the line, the data on the short sections of isolated pipe are of interest, although, for reasons already explained, there are not many

of these data.

If all the coated pipe that developed measurable pits during both the 4-year and 10-year periods of exposure are considered, there are 18 cases out of a possible 107 in which the rate of penetration was higher for the longer period of exposure. In other words, about 17 percent of the specimens to which coatings had been applied showed an increasing rate of pitting. There were 35 cases out of a possible 67 in which the quotient obtained by dividing the maximum pit depth on the coated pipe by the average maximum pit depth on the uncoated pipes at the same site was greater for the 10-year period of exposure than for the 4-year period. This is equivalent to saying that for 52 percent of the pipe protected by coatings, the maximum rate of penetration decreased more slowly than on bare pipe at the same site. This comparison is somewhat too favorable to the coatings, as no account has been taken of the pipe protected by coatings that were punctured at the close of the 10-year period or of the cases where the coated pipe showed no measurable pits at the end of the 4-year period, but did show measurable pits at the end of the longer period of exposure. In a general way at least, the data for the coatings on the line are in agreement with those for the coatings on the isolated pipe. Although it is possible that the cases which indicate that the coatings accelerated the corrosion are accidental, the percentage of such cases makes this seem doubtful. The implication of the data that have been presented raises a serious question as to the ultimate usefulness of coatings which are imperfectly applied, contain pinholes, or allow the pipe to become exposed because of pipe movements, soil stress, penetration by roots, or deterioration of the coating materials.

The available data are insufficient to justify positive and final conclusions as to the explanations for the shapes of the curves that have been presented or as to their significance. Nevertheless, the data are sufficient to warrant very thoughtful consideration, as they may contain an explanation for much of the pipe-line trouble that has been

experienced.

The reader should of course keep clearly in mind the fact that even an accelerated rate of pitting on coated pipes which are not completely protected does not imply that the coating will not prolong the useful life of the pipe, as occasional leaks may be repaired at no great cost. The summary of the data in the final report [61] indicates quite clearly that even the least effective coating reduced the number of feet of pitted pipe and that less than 2 percent of the footage of the pipe with the most effective coating showed any pits. The reduction in the total number and average depth of pits on the coated pipe was much greater than is shown by the table referred to.

The solution of the problem of pipe-line protection seems to be better coatings supplemented by cathodic protection to take care of the imperfections that are not avoided by better materials and better application.

(6) Conclusions based on the API Tests.—The data on the conductance of the coatings lack uniformity. Not infrequently one section of a coating showed many times the conductance of another section of the same coating at the same test site. In many cases it is difficult or impossible to determine whether this lack of consistency is the result of lack of uniformity in the thickness or composition of the coating, poor application, or of nonuniform soil conditions. The same statements are applicable to maximum pit depths. The important fact is that the amount of protection afforded by many coatings was not uniform even under nominally the same condition. Either the coatings as applied were not uniform or they lacked a sufficient factor of safety to provide against the variations in the exposure to which they were subjected.

There was a general increase in the conductance of the coatings over that observed in 1934. This indicates that the usefulness of the coatings is limited with respect to time. There are individual exceptions to this

generalization.

As a class, the coatings that were applied cold and were not reinforced or shielded showed more cases of exposed pipe than did the thicker

coatings

Zinc chromate and baked-on red and blue lead appear to be satisfactory as undercoats under the conditions to which they were subjected. The bond between the undercoats and the pipe was stronger than that between the undercoats and the coatings applied over them. These undercoats seem to have reduced corrosion, but did not afford adequate protection to pipe in corrosive soils.

The number of coated pipe showing patterns was greater in 1940 than in 1934, although some sections that showed a positive pattern in

1934 showed none in 1940. The pattern tests indicate a continued decrease in the effectiveness of most coatings.

Only four test sites were severely corrosive, as indicated by the depths of the pits on unprotected sections of pipe. The majority of the test sites were, therefore, favorable to the coatings, if pit depths are used as criteria for the effectiveness of the coatings. There is no consistent difference between the depths of the deepest pits on uncoated sections of line pipe and on the uncoated short pipe sections.

On the average, the pipe under more of the coatings showed corrosion in 1940 than in 1934, and the maximum pit depths were greater for

most coated sections.

In general, there were fewer pits under thick coatings than under thin ones, but the depths of the pits under a thick coating were sometimes

greater than under a thinner one.

Because the coatings differ in many ways, it is impossible to determine positively whether the machine-applied coatings were superior to those applied by hand or to determine the relative merits of coal tar and asphalt as protective coatings. Likewise, the data do not show clearly whether impregnated asbestos felt was superior to organic materials as a reinforcement. However, as a shield it did not deteriorate as did the organic materials. A layer of cement-sand mortar over an emulsion coating was insufficient to prevent serious pitting in a corrosive soil.

All the coatings reduced the number of pits per unit area of surface, and most of the coated sections of pipe showed shallower maximum pit depths than occurred on the corresponding unprotected pipe. Several of the coatings afforded nearly complete protection to most of the sections of pipe to which they were applied, but no coating completely

protected all sections.

The primary purpose of the test was to establish engineering principles related to the manufacture and use of protective coatings for pipe lines. The following are some of the facts established by the test.

1. Many of these coatings will greatly reduce corrosion during at least 10 years (the period of this test), although complete protection from all corrosion has not been realized in corrosive and destructive soils.

- 2. The effectiveness of all coatings tested decreased throughout the period of the test. This in most cases is the result of continued soil pressure and the absorption of water. There appears to be little change in the coating materials other than that in the organic fabrics used as reinforcements or shields.
- 3. Shields and reinforcements should be permanent and sufficiently rigid to distribute soil stress and pressure due to the weight of the pipe over enough area to prevent the flow of the bituminous or other material used in the coating.

4. Thickness of material is an important factor in coating construction.

- 5. A coating should be sufficiently rigid to withstand pressures over long periods and elastic enough to withstand stresses resulting from pipe movement and sudden changes in temperature. These requirements are difficult to reconcile.
- 6. Under certain conditions, a coating that fails to protect at a few points may cause deeper pits than would have occurred on uncoated pipe in the same location.
- 7. Many soils are only mildly corrosive, and no protective coating is required in them unless the cost of a leak would be abnormally high.

8. The economics of the general problem should be considered.

### 3. NONBITUMINOUS, NONMETALLIC COATINGS

Among the corrosion-resistant materials buried by the National Bureau of Standards in 1932, and subsequently, were a number of specimens of newly developed nonbituminous, nonmetallic coatings, some of which had been applied to short sections of pipe lines. Others have not vet been sufficiently developed to justify such trials. The Bureau's tests should be regarded as preliminary or elimination tests, as they are subject to the limitations already discussed. It is possible that, at present, some of the coatings under observation are too expensive for general use. In a number of cases the problems of field repairs that are unavoidable and the application of the coatings to joints in the line and to fittings have not been solved. Nevertheless, the results of the tests are of interest and suggest new ways of attacking the pipe-protection problem. The coatings are described in table 12.

#### (a) AFTER 9 YEARS OF EXPOSURE

Table 82 shows the condition of the pipe beneath the coatings exposed for 9 years. It will be noted that the two thick rubber coatings completely protected the pipes except at the ends where moisture got be-

Table 82 .- Condition beneath nonmetallic coatings exposed for 9 years. [Figures indicate depths of maximum pits in mils]

E, ends corroded. G, ends rusted. H, pipe punctured, 145 + mils.
M, metal attack—pipe roughened by corrosion. R, pipe rusted.
P, definite pitting but no pits greater than 6 mils.
U, no corrosion.
D, pipe destroyed by corrosion.

	Soil	Specimen No.	Porcelain enamel	Rubber paint	Paint	Cashew-nut-oil- asbestos fiber	Hard rubber	Loaded hard rubber	Synthetic resin	Chlorinated rub- ber paint	China wood-oil mica compound	Uncoated low-carbon- steel pipe
No.	Type		С	D	E	F	G	н	J	K	w	N
53 55 56 59 60 63 64 65 66	Cecil clay loam Hagerstown loam Lake Charles clay Muck Carlisle muck Rifie peat Sharkey clay Susquehanna clay Tidal marsh Docas clay Chino silt loam Mohave fine gravelly loam Cinders	123121212121212121 21 21 21 21	DE: RUDE E E E E DE DE 43E	M 9 MM *511 *383 522 199 MM 155 *44 RR UU 233 M 37 47 121 129 15 777 H	17 15 16 18 197 197 197 197 15 138 25 25 25 25 10(7) 287 77 122 D(7) H	MR :RRMM61RMR31RRRRR27P :M MPR34 M66 R	ממממי	 U U U	M M		MM MMMMMMM GG MM MM GG MM MM GG MM MR ME	59 59 59 

^{*} Severe uniform corrosion; no reference surface; impossible to measure true penetration.

b The number in parentheses after the pit depth indicates that 1 or both specimens was punctured in the previous removal indicated by the number, (5) indicates that the pipe was punctured after 5 years, etc.

neath the coating, as it might also at points where the coating on a pipe is injured. The porcelain enamel was nearly as good. It is probable that some of the corrosion reported was the result of injuries in handling the specimens. However, as much greater care was taken than is practical with coatings on pipe lines, the corrosion resulting from injuries should not be entirely disregarded. The thick china-wood oil compound, W. prevented serious pitting in all soils. The fact that many of these coatings cracked and had a strong odor, and the blistering of the priming coat indicate that the coating had deteriorated. However, because of its thickness, it may remain helpful by separating the pipe from the soil for several more years. The thin coatings (D, E, F, J, and K) definitely deteriorated and are probably of little further use. Most of these coatings showed evidence of breakdown on earlier examinations.

#### (b) AFTER 4 YEARS OF EXPOSURE

Four of the five coatings buried in 1937 were made from Bakelite. The other was Koroseal. Tables 83 and 84 show the condition of the pipes beneath the coatings. Two coatings remained unaffected at one

Table 83.—Condition of metal beneath Bakelite and Koroseal coating exposed for 4 years.

U = No corrosion.

□ = No corroson.
 R = Pipe rusted.
 M = Metal attack—pipe roughened by corrosion.
 P = Definite pitting but no pits greater than 6 mils.
 E = Links corrodon.

			C	oating symb	ool		Uncoated low-carbon-
Soil No.	Specimen No.		Bakeli	te base		Koroseal	steel pipe exposed 5 years,
		L	M ₁	M ₂	N	P	N N
53	$\left\{ \begin{array}{cc} 1\\2 \end{array} \right.$	U R	R R	M M	R R		47 53
55	$\left\{ \begin{array}{cc} 1 \\ 2 \end{array} \right.$	U 18	P P	M M	R R		48 66
56	$\left\{ egin{array}{c} 1 \ 2 \end{array}  ight.$	R R	P P	58 M	P P		53 88
58	$\left\{ \begin{array}{cc} 1\\2 \end{array} \right.$	U 12	P 15	ME 25	P R		97 108
60	$\left\{ \begin{array}{cc} 1\\2 \end{array} \right.$	R U	P 12	55 PE	M R		18 29
61	$\left\{ \begin{array}{cc} 1\\2 \end{array} \right.$	18 U	25 R	25 18	R R		23 85
62	$\left\{ \begin{array}{c} 1 \\ 2 \end{array} \right.$	ភ្ជ	R M	M M	R R	15 15	66 66
33	$\left\{ \begin{array}{c} 1\\2 \end{array} \right.$	R R	M R	M M	10 13	::::::::	33 38
34	$\left\{ \begin{array}{cc} 1\\2 \end{array} \right.$	R R	r L	P P	U R		154+ 164+
35	$\left\{ \begin{array}{c} 1\\2 \end{array} \right.$	R. U	R R	13 M	R M		79 69
36	$\left\{\begin{array}{cc} 1\\2\end{array}\right]$	M 22	R P	12 28	M P	12 P	154+ 154+
57	$\left\{\begin{array}{cc} 1\\2\end{array}\right]$	R R	120 107	41E 57E	35 30	45 50	84 154+
9	$\left\{ \begin{array}{c} 1 \\ 2 \end{array} \right]$	39 M	24 25	R 32	M 15	P 8	
70	$\left\{\begin{array}{cc} 1\\2 \end{array}\right]$	U R	R R	PE 16	P P		

test site. Corrosion had started beneath all the other specimens. The failures of the Bakelite coatings were usually accompanied by blistering. The failure of the Koroseal coating is characterized by a deterioration of the fabric to which Koroseal is applied.

Table 84 .- Summary of data on condition of coated pipe exposed for 4 years.

[Figures in the table refer to the number of test sites. Data from specimens buried in 14 test sites except for P specimens, which were buried in 4 test sites. Poorer of the 2 specimens reported.]

		Condition	of pipe		Coating
Coating symbol	Unaffected	Rusted	Roughened	Pitted	blistered
L. M 1. M 2. N 2. P a	1 0 0 0	8 3 0 5	0 2 4 2 0	5 8 10 7 4	6 13 14

a Poor bond between pipe and coating.

#### (c) AFTER 2 YEARS OF EXPOSURE

In 1939 two types of coatings were buried at 14 of the test sites, and an additional coating was placed at four of the sites. See table 12 for the maker's description of these coatings. Two specimens of each coat-

Table 85.—Condition of coated pipe exposed for 2 years.

t = No corrosion.					
Soil No.	Specimen No.	Blue basic lead sulfates phenolic resin varnish paint,	Plastic- treated fiber,	Thiokol,	Uncoated low-carbon- steel pipe
		R	s	т	N
53	$\left\{ egin{array}{c} 1 \ 2 \end{array}  ight.$	M M	P P		37 37
55	$\left\{ egin{array}{c} 1 \ 2 \end{array}  ight.$	M M	P P		40 42
56	$\left\{ egin{array}{c} 1 \ 2 \end{array}  ight.$	12 M	25 20		26 13
58	$\left\{ egin{array}{c} 1 \ 2 \end{array}  ight.$	M M	P P	R R	20 16
60	$\left\{ \begin{array}{cc} 1\\2 \end{array} \right.$	R R	M M		55 19
61	$\left\{ egin{array}{c} rac{1}{2} \end{array}  ight.$	M M	M M	R. U	8 13
62	$\left\{ egin{array}{c} 1 \ 2 \end{array}  ight.$	M M	10 10	R R	68 36
63	$\left\{ egin{array}{c} 1 \ 2 \end{array}  ight.$	M M	28		21 9
64	$\left\{ egin{array}{c} 1 \ 2 \end{array}  ight.$	$_{ m M}^{ m R}$	P 20	R R	123 138
65	$\left\{ \begin{array}{cc} 1 \\ 2 \end{array} \right.$	$_{\mathbf{M}}^{\mathbf{M}}$	P P		43 36
66	$\left\{ \begin{array}{cc} 1\\2 \end{array} \right.$	55 P	8 12		71 61
67	$\left\{\begin{array}{cc} 1\\2\end{array}\right.$	M 27	27 29		154+ 154+
69	$\left\{ \begin{array}{cc} 1\\2 \end{array} \right.$	M M	M M		
70	$\left\{ egin{array}{c} 1 \ 2 \end{array}  ight.$	10 18	8 8		

M = Metal attack—pipe roughened by corrosion. R = Pipe rusted. P = Definite pitting but no pits greater than 6 mils. U = No corrosion.

ing were removed from each test site in 1941. Table 85 shows the condition of each coated pipe after an exposure of 2 years. Only one speci-

men showed no sign of corrosion.

The lead-sulfate coating, R, blistered and became scaly. There was little adhesion between the plastic coating S and the pipe. The entire surface of each pipe coated with the plastic rusted. Soil indented the coating and adhered to it. In most soils the coating appeared to have rotted.

On each specimen of Thiokol-coated pipe there was a network of fine cracks which usually did not permit water to reach the pipe. In a number of cases the coating was chipped, possibly by the formation of rust beneath the coating.

# IX. LABORATORY STUDIES OF CORROSION 1. DENISON CORROSION CELL

Denison [80, 81] has developed a corrosion cell for the study of corrosion phenomena in the laboratory. With this cell, the behavior of various soils and metals can be investigated under uniform conditions of moisture, aeration, etc., and the current through the cell can be regulated at will.

The cell consists essentially of two electrodes of the same metal separated by a layer of moist soil, which constitutes the electrolyte. Differential aeration of the two electrodes is brought about by making one of them more accessible to the air than the other, whereby the cell develops an internal electromotive force. It is often desirable, however, to control the current through the cell, which is done by means of an external battery and variable resistance. The construction of the cell is described in detail in Appendix 5. In the use of the Denison cell for investigating corrosion phenomena, the principal measurements made are of opencircuit potentials, short-circuit currents and simultaneous values of current and potential under various conditions.

Open-circuit potentials are measured with a potentiometer-voltmeter, which has the advantage that it draws no current from the circuit under test. Individual potentials of the anode and cathode are determined by using an auxiliary saturated calomel electrode connected through an agar-KCl bridge placed in contact with the soil through a hole in the

bottom of the cell shown in figure 87 in Appendix 5.

Short-circuit currents are measured by means of a "zero-resistance" milliammeter in which the resistance of the instrument is compensated for by opposing the voltage drop in the instrument by an equal applied

voltage.

The measurement of potentials in the cell while current is flowing is complicated by IR drops in the circuit. If the resistances were constant and accurately known, the IR drops could be calculated and proper corrections could be made, or, under certain conditions, errors from this source could be made small by placing the tip of the reference electrode close to the electrode whose potential is being measured, or by placing the reference electrode out of the path of current flow. According to Hickling [82], however, the error resulting from inclusion of some IR drops in the measured potential is seldom avoided, as there always is present a certain surface resistance which may be due to gas evolution, depletion of electrolyte in the layer in contact with the electrode, or, in the case of the anode, to a poorly conducting film. Wholly apart from the error due to the resistance of films, the high resistivity of soils

renders the measurement of potentials particularly difficult. Furthermore, marked variation of resistance with current may occur. The above methods are not readily applicable to the Denison cell.

Errors due to these unknown and variable IR drops can be practically eliminated by using a method developed by Hickling [82] and adapted to the Denison cell by Darnielle [83]. In this method, the current is periodically interrupted for very short intervals of time, and the potentials are measured during the period of interruption. If the interval of interruption is too long, the potential of a polarized electrode will change during the measurement, so that the observed value will not correspond to the potential existing while the current is flowing. The advantage of the Hickling method is that the time during which the current is interrupted can be made very short (of the order of 10⁻⁵ seconds), so that errors due to depolarization are very small. Details of the method are given in Appendix 6.

From data obtained with the corrosion cell, various relationships and critical values can be determined which appear to be related to the type of control, the rate of corrosion, the relative corrosiveness of different soils, and the relative resistance to corrosion of different metals. The principal relationships which have been useful in this connection are represented by current density-potential curves and potential-time curves.

The rate of corrosion of steel under many soil conditions appears to be controlled chiefly by the rate at which the cathodic areas are depolarized. In fact, in the absence of conditions which favor the formation and development of protective anodic films or of corrosion products in immediate contact with the corroding area, the progress of corrosion in soils is probably determined by the rate of the cathode reaction. Corrosion under such conditions is said to be under cathodic control. On the other hand, if a more or less protective film develops on the anode, the rate of corrosion tends to be controlled by the reaction at the anode and the corrosion is then said to be under anodic control. The type of control under which corrosion proceeds, that is, whether anodic or cathodic has been shown by Bannister and Evans [84], Burns [85], and Brown and Mears [86] to be indicated by the relation between the current (or current density) and the potential of the anode and cathode of a corroding specimen. Current density-potential curves that are typical of cathodic control are shown in figure 42. It will be seen that the change in the cathode potential with increasing current density is much greater than the corresponding change in anode potential. This is particularly well illustrated by the curve for soil 57. A current density of 17 ma/dm² raised the potential of the anode by only 0.05 volt, whereas the potential of the cathode was lowered by 0.20 volt. Curves for metals corroding under anodic control show that the change in potential with increasing current density is greater for the anode than for the cathode.

If it were not for the high internal resistance of the cell, which limits the value of the current on short circuit, the rate of corrosion could be determined simply by measuring the short-circuit current at a suitable stage of corrosion. It is preferable, however, to eliminate the effect of resistance. This is accomplished by making use of the current density potential curves. As the current density is increased, the potentials of the anode and cathode approach the same value. At the point where the two curves intersect, the cell is completely polarized, the internal potential difference is zero and the applied electromotive force just balances

the internal IR drop. The potential of the electrodes at this point is known as the corrosion potential. The associated current density is the limiting value for the particular combination of soil and electrodes and corresponds to the current for the maximum rate of corrosion. The interpretation of current-density-potential curves as indicative of rate of corrosion has been discussed by Evans [87], Evans and Hoar [88], Burns [85], and others.

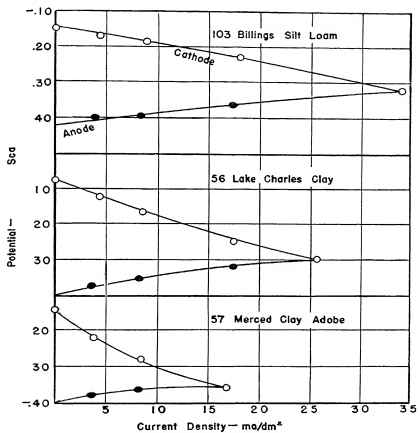


FIGURE 42.—Current-density-potential curves.

Metals such as stainless steel, lead, copper and copper alloys, which tend to form protective oxide films in soils, develop only small potential differences and as such a cell has a high resistance only a negligible current flows between the electrodes. Under these conditions, the corrosion is due almost entirely to local cell currents and obviously the short-circuit current would not be sufficient to account for the observed corrosion. If it were not for local action, it should be possible to stop corrosion at the anode by opposing to the potential difference developed internally

an external electromotive force just sufficient to prevent current from flowing away from the anode. However, in order to completely suppress the local action, which is the principal cause of corrosion in the cell when on open circuit, it is necessary to apply a greater electromotive force so that current flows into the anode. The minimum value of current required to protect the electrode from corrosion is called the protective current.

The local-cell theory of Müller [89] indicated that, with cathodic control, the protective current should be equal to the corrosion current. Other investigators have shown that the protective current is equal to the corrosion current under certain conditions. For example, Gatty and Spooner [90] found good agreement between values which they obtained for the "critical," that is, protective, current for copper in dilute sulfuric acid and in neutral salt solutions and the currents calculated from data available in the literature on corrosion under similar conditions. According to Mears [91] the amount of current required to prevent corrosion is equal to the corrosion current only if the corrosion reaction is completely determined by oxygen diffusion, that is, by cathodic control.

Evans, Bannister, and Britton [92], working on the current-potential relation of various metals when made cathodic in an aerated solution of 0.1 N potassium chloride, found that, at current densities below the minimum value required to protect the metals cathodically, the potential of the cathode remained unchanged as the current density was increased. At current densities greater than the protective value, the potential of the cathode increased logarithmically. This behavior may be explained by assuming that the plate was originally made up of anodic and cathodic areas and that the latter were covered by a high-resistance film. Current therefore flowed only to the anodic areas until they were destroyed by the neutralizing of the local currents, after which the entire plate becomes cathodic. Ewing [93] has utilized this principle for determining the protective current by plotting the potential of the cathode against the logarithm of the applied current.

#### 2. BEHAVIOR OF STEEL IN VARIOUS SOILS

Denison and Darnielle [94], using the corrosion cell, have investigated the behavior of steel in various soils when it corroded under cathodic control. Measurements of the corrosion potential were made at intervals during a period of 2 weeks. During the test period, the cells were kept on short-circuit and maintained at a temperature of 25°±0.5°C. The moisture content of the soils was maintained constant by placing the cells in separate friction-top cans in which a small quantity of water was placed in such a way that it did not come into direct contact with the cells.

Typical corrosion-potential—time curves are shown in figure 43. It will be observed that after the first few days the corrosion potential became practically constant. In a few soils, a slight regular increase in the corrosion potential was noted even at the end of the test period. This is attributed chiefly to the fact that the potential of the cathode tends to become more positive with time. In all the soils studied, the corrosion potential was more positive than the open-circuit potential of the anode. Brown and Mears [86] have pointed out that the potential of a galvanic couple can be identical with the open-circuit potential of the anode only if there is no anodic polarization. The fact that in many soils the polarization curve for the cathode is very flat causes the potential of the short-circuited electrodes to become more cathodic.

Typical results of measurements of open-circuit potentials of the anode and cathode over a test period of 2 weeks are shown in figure 44. The marked change in the potential of the cathode (which becomes more positive) indicates that the potential of that electrode tends to approach the potential of the oxygen electrode in the particular environment.

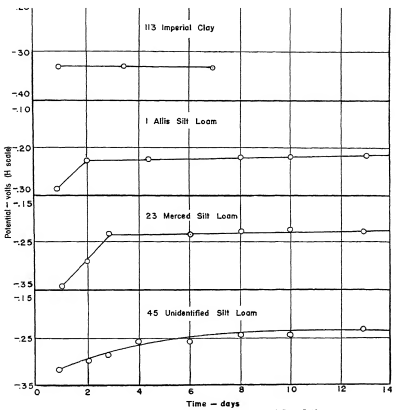


FIGURE 43.—Relation between corrosion potential and time.

The reason the latter potential is not reached is that imperfections in the protective oxide film expose the underlying metal. During the initial period of exposure, anodic areas develop on the perforated electrode as well as on the solid disk. As the solid disk is less accessible to air, the areas surrounding the points of attack on the disk soon become cathodically polarized. As a consequence, current leaving the anodic area on the solid disk flows to the perforated disk, where, because of the relatively greater concentration of oxygen, cathodic polarization is less. The extension of the cathodic areas on the perforated disk diminishes the number of anodic areas because the cathodically produced OH ions in the presence of oxygen precipitate ferric hydroxide in immediate contact with the corroding areas and thereby effect repair of the film.

In addition to the measurement of potential, daily observations of the short-circuit current of the cells were made. The maximum current developed usually within 18 to 24 hours, although in a few cells the current increased gradually for a week or longer.

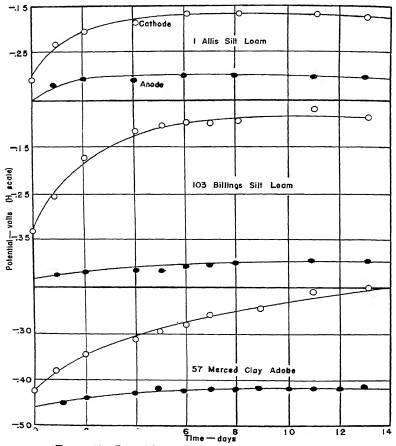


FIGURE 44.—Potential-time curves for the anode and cathode.

In order to correlate the results of the various electrical measurements with the corrosion, the loss of weight of the anodes was determined. The results are summarized in table 86, in which the soils are arranged in the descending order of their corrosiveness as indicated by the loss of weight during the 2-week test period. It can be seen that there is a rather definite correlation between the loss of weight and either the current at corrosion potential or the maximum short-circuit current. The ratio between these two currents is fairly constant in soils of low resistivity, and tends to increase in soils of higher resistivity, as would be

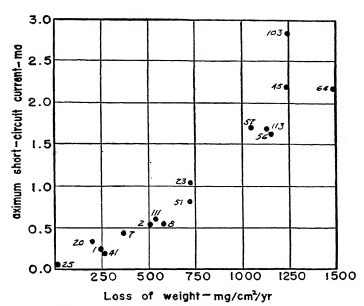


Figure 45.—Correlation between maximum short-circuit current and loss of weight of the anodes.

expected. The correlation between the maximum short-circuit current and loss of weight is shown in figure 45. The somewhat erratic nature of the results can be ascribed to the unequal ratio of the maximum to the average current for the different soils.

Table 86.—Corrosiveness of soils as indicated by electrical measurements and loss of weight of the anodes.

Soil No.	Soil type	Maximum open circuit voltage ^a	Corrosion potential b	Current at corrosion potential	Maximum short- curcuit current	Loss of weight c
64 103 45 56	Billings silt loam Unidentified alkali soil Lake Charles clay	0.31 .30 .38 .37 .38	0.34 .32 .29 .29 .31	ma 2.72 3.86 2.55 2.92 2.90	ma 2.19 2.85 2.20 1.64 1.70	(mg/cm²);yr 1,475 1,230 1,230 1,150 1,130
57 23 51 8	Acadia clay	.30 .34 .12 .12 .16	.35 .22 .31 .37 .21	2.00 1 88 1.20 0.78 1.20	1.70 1.05 0.80 .55 .60	1,040 722 708 580 534
2 7 1 41 20	Sur ni. srt loam		.33 .24 .22 .26 .30	0.90 .50 .60 .40 .67	.54 .42 .23 .20 .32	504 370 244 214 202
25	Miami clay loam	.01	.32	.005	.003	16

[·] Potential between anode and cathode of corrosion cell on open circuit.

Potential at intersection of current-density-potential curves.
 Calculated from results in a 2-week run.

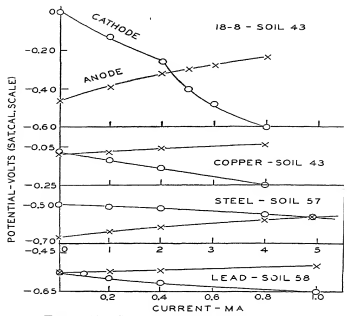


FIGURE 46.—Corrosion under cathodic control.

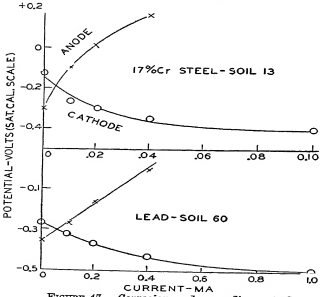


FIGURE 47.—Corrosion under anodic control.

### 3. BEHAVIOR OF DIFFERENT METALS IN VARIOUS SOILS

In view of the extensive use of metals other than iron or steel, such as copper, brass, lead, zinc, etc., Denison [95] has used his corrosion cell for the investigation of the behavior of several different metals. In most soils, the corrosion of steel is under cathodic control. However, if a protective film develops on the anode, the corrosion tends to be anodically controlled. The type of control for the various metals in different soils was determined by inspection of the current-potential curves of the separate electrodes. The typical curves shown in figures 46 and 47 indicate corrosion under cathodic and anodic control, respectively.

It is of interest to consider the behavior of lead, as illustrated by the current-potential curves for lead in figures 46 and 47, with respect to the environmental conditions of Houghton muck (soil 58) and Rifle peat (soil 60). Both soils are organic, having been derived from the decomposition of vegetable matter. Reference to the composition of the soluble material in the soils (table 20) shows that both soils are very acid and that soil 60 is extremely high in sulfates. The slight anodic polarization shown by lead in soil 58 (figure 46) is consistent with the known corrosive action on lead of organic acids produced in the decomposition of wood and other organic matter. The organic acids maintain a low concentration of lead ions at the anode. The corrosion of lead in soil 60 was inhibited by the marked polarizing action of sulfate ions, exhibited

by the curve for the anode in this soil (figure 47).

The curves for stainless steel of the 18-8 variety in soils 43 and 13 represent the behavior of this alloy in the active state (figure 46) and in the passive state (figure 47). While in the passive state, even a slight flow of current is seen to have a marked effect on the potential of the anode. However, with the metal in the active state, the anode exhibits the characteristic behavior of ordinary carbon steel. The shift from the passive to the active condition in soil 43 resulted from the inability of the oxide film to maintain a state of repair in a reducing environment high in chlorides and sulfates. In figure 46 the change in curvature for the cathode at 2 milliamperes corresponds to the shift from the oxygen electrode reaction to that of the hydrogen electrode. Curves of this type

have been discussed by Hoar [96].

In table 87 is shown the type of control under which the materials corrode in the different soil environments. The corrosion of zinc and steel is under cathodic control in all of the soils because, in the presence of chloride, sulfate, carbonate, and bicarbonate ions in an environment deficient in oxygen, the primary reaction products of steel and zinc are soluble and consequently diffuse readily away from the anode surface. Under these conditions, a fairly large negative potential can be maintained at the anode, even at relatively high current densities. However, in very porous soils which are either strongly alkaline or are deficient in soluble salts, the type of control of steel and zinc shifts from cathodic to anodic. The fact that the corrosion of zinc is generally under cathodic control has naturally an important bearing on its use as a protective coating for steel and as an anode in cathodic protection installations.

Following the procedure of Ewing [93] for determining protective current, the potential of the cathode was plotted against the logarithm of the applied current for a variety of metals in Docas clay (soil 64). The curves are shown in figure 48. Most of the curves are composed of two parts, the first being horizontal (or nearly so) and the second being linear with a negative slope. At the higher currents, the cathode is polarized to such values that hydrogen evolution begins and the potential

follows the linear hydrogen overvoltage curve. Similar curves have been obtained and discussed by Gatty and Spooner [90] and by Hoar [96]. In the case of a few metals in certain soils, the first part of the curve has an appreciable negative slope. This change in potential may have resulted from a partition of the applied current such that a certain fraction flowed to the cathodic areas. Under such conditions, which would be caused by the high resistance of small anodic areas, the protective current would be greater than that associated with corrosion.

Table 87 .- Control of the corrosion rate of metals in different soil environments.

				Envi	ronment			
Material	Alkali car	bonate	Chloride	Sulfate	Chloride and	Reducing, with	Reducing, with organic	Reducing with organic acids chloride
	Good aeration	Poor aeration			sulfate	organie acids	acids and sulfate	and sulfate
			S	oil number				
	13	a 23	64	57	56	58	60	43
Low-carbon steel	Cathodic b	Cathodic	Cathodic	Cathodic	Cathodic	Cathodic	Cathodic	Cathodic
Steel with 17% Cr	Anodic	Anodic	Anodic	Anodic	Anodic	Anodic	Anodic	do
18-8 steel	do	do	do	do	do	do	do	do
Copper	do	do	do	do.,	do	Cathodic	Cathodic	do
Brass (60-40) .	do	do	do	do	do	do	do	do
Zine	Cathodic	Cathodic	Cathodic	Cathodic	Cathodic	do	do	do
Lead	Anodie	Anodic	Anodic	Anodic	Anodic	do	Anodie	Anodic

#### 4. COMPARISON OF LABORATORY DATA WITH THE RESULTS OF FIELD TESTS

In view of the fact that conditions in the field are not uniform, an exact correlation between the results of laboratory tests and those obtained in the field should not be expected. The laboratory tests, however, should be expected to give a general indication as to whether under given field conditions corrosion will be severe or negligible. In order to determine to what extent this is true, the results of the laboratory tests have been compared with those obtained in the field for the same metals buried in the same soils. The corresponding values are given in table 88. In order that the laboratory data may be made comparable for the different materials, the loss of weight corresponding to the selected value of corrosion current or protective current was calculated. The values thus obtained can be compared with the actual losses during the exposure period of 2 weeks in the case of the cells that had a definite polarity. As the cell current was necessarily less than the current at corrosion potential, on account of the IR drop in the cell, the calculated losses are greater than the actual losses, except in cells of very low resistance.

Generally speaking, the current at corrosion potential can be taken as the measure of the rate of corrosion whenever corrosion proceeds under cathodic control. An exception must be made, however, for two organic soils, Houghton muck (soil 58) and Rifle peat (soil 60). On account of

^{*} Sulfates also present.

b Anodic with exceptionally good aeration.

the very high water content on these soils, which approached saturation. the perforated-disk cathode could not develop a potential sufficiently noble to overcome the resistance of the soil. Whenever comparisons are possible, the agreement between the values of protective current and the current at corrosion potential is generally good, which indicates that protective current can also be taken as a measure of corrosion rate, at least when the rate is controlled cathodically. The few cases of poor agreement can be reasonably accounted for.

The field data given in table 88 show the average rate of loss of weight and rate of maximum penetration, usually of two specimens of each material after exposure to the various soils for approximately 8 years.

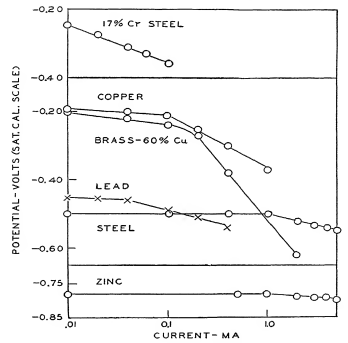


FIGURE 48 .- Current-potential curves for the cathode in Docas clay.

When data for more than one grade of a given material were available, as was the case with lead and zinc, the values for all grades were averaged. Because of the marked tendency for the rate of corrosion to decrease with time of exposure under certain environmental conditions, the rates given for lead and zinc after exposure for 2 years are not strictly comparable with the values given for the usual 8-year period.

To facilitate the comparison between the field and laboratory data, the relative order of the metals with respect to their resistance to corrosion is given in the table. The correlation of the laboratory data with loss of weight in the field is somewhat better than with the rate of pitting, which seems reasonable in view of the fact that a given loss in

TABLE 88.—Comparison of corrosion data from Jaboratory and field tests.

		2 20 10 10 10 10 10 10 10 10 10 10 10 10 10		-	לי מיני לי מיני מיני מיני לי מיני מיני מ		and annua for	10 01 000.		To the second se	
			Laborat	Laboratory data			Field data		Order of ex	Order of corrodibility based on-	ased on-
Material	Type of control: C=cathodic A=anodic	Current at corrosion potential	Protective	Calculated loss in weight	Actual loss in weight	Арргохі- пясь ехромиге	Rate of loss in weight a	Rate of pitting	Calculated loss of weight in laboratory test	Rate of loss of weight field test	Rate of pitting in field test
			SOIL	13, HANFO]	SOIL 13, HANFORD FINE SANDY LOAM	INDY LOAD	A.				
Steel Zine Copper Brass (60-40) Lend 17% Cr steel	DDAAAA	ma 0.70 1.80	ma 0.50 30 28 32 32 .06	0.245 .738 .111 .127	0.137	<u>2</u> 0000000	(oz/ft³)/yr 1 24 0.31 .03 .10	Mils/yr 11.8 6.3 1.0 1.0	ಬ≎ಐೆೆೆೆೆೆೆೆ	5248314	ರಾಣ ಅಹ 🗆 🛏
			80]	IL 23 (70), 1	SOIL 23 (70), MERCED SILT LOAM	LT LOAM					
Steel Zine. Copper Brass (60-40). I.ead	OD4444	2.40	1.30 .70 .03 .06	0.455 .984 .012 .024 0	0.457	∞α∞∞ •	2.45 0.74 0.75 .03	19.6 20.0 1.9 0	70 00 4 HH	0 4 8 70 8 T	7.040 C
				SOIL 43,	TIDAL MARSH	RSH					
Steel.	೦೧೧೬೦೧೧೧	1.00 3.60 0.40 0.18 3.40	1.20 0.42 .16 .08 .11	0.350 1.476 0.167 .064 .103 1.190 0.770	0.246 .927 .114 .010 .106 .462	&&&&&&	1.40 0.21 .60 .03 .05 .67	8.1 8.1 (d) 1 30.0 1	41-10-101010	F4100800H	9 5 5 7 4 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7
			SS	OIL 56, LAP	SOIL 56, LAKE CHARLES CLAY	S CLAY					
Steel	೦೦44444	2.20	0.90 0.035 0.035 0.04	0.326 .902 .014 .016 .129 0	0.334	7. 2. 5 2. 5 7. 5 7. 5	2.75 0.84 0.88 .09 .15	16.5+ 9.0 1.1 1.0 24.5	07:84:01L	r-@8342519	° 0 20 20 20 20 40

9745887

ಹಿ 4 ಬ ಟ ⊣ ಗು

0407000-

7 2 3 4 T 10 C)

r48=01031

アちょうけいこ

	F084111		೦೯೮೮೩೩		\$2500 # H 450 H		<b>664684</b> 4	
	19 0 1.1 4.2 4.2 1.0		14.4 14.1 1.8 1.1 10.7		32.5 31.8 1.2 10.1 1.1		1880001 100000+1 100000+1	
	1.66 0.02 30 .018 .0002		1 2 76 1 0 54 2 22 6 1 27 1 27		1.45 3.03 0.14 .07 .045		4 88 0 51 0 13 0 10 0 00 0 00 0 00 0 00 0 00 0 0	
ADOBE	வவ வவ வ		88.00 8.00 8.00 8.00	ΑT	1181181	AY	741074107	
SOIL 57, MERCED CLAY ADOBE	0 627	SOIL 58, MUCK		SOIL 60, RIFLE PEAT		SOIL 64, DOCAS CLAY	0.627	
IL 57, ME	1 715 0 964 .032 .028 0 0	NOS	0.455 .160 .044 .076 .129	SOIL 60	0.290 .328 .111 .111 .135 .065 .065	BOIL 04	0.770 2.870 0.040 0.040 0.039 0.039	
SOS	2.15 0.08 0 0 0		1.60 0.25 0.25 11 .19 .10		0 88 82 83 83 83 83 83 83 83 83 83 83 83 83 83		1.8 0.10 .18 .03 .005	.05.
	4.02 8.83 7.53		1 30 0 33				7.5	, multiply by 3.
	೮೮४४४४४		೯೦೦೦೦೦		DDDD444		<b>೮</b> ೦ <b>४४४४४</b>	g/dm²)/yr
	Steel. Zine. Brass (60-40). 17% Cr steel.	and the second s	Steel. Zine. Zine. Bruss (00-40). Lead. 17% Ct steel.		Steel. Zino. Zino. Brass (60-40) 17% Ct steel.		Skeel. Zino. Zino. Brass (60-40). Irad C. skeel. 18-8. Cr skeel.	To convert these values to (g/dm²)/yr, multiply by 3.05.

* To convert these values to (g/dm?) fyr, multipuy by .b Yalue sammet, to faellitate comparison of data.

• Data for steel with 20% Cr.

• Data for stringer soil 63.

• Data for similar soil 63.

weight might be distributed fairly uniformly over the surface, as was the case with copper in soil 43, or it might be confined to a relatively

few pits, as in the case of the stainless steels.

Both the laboratory tests and the field data indicate that the corrosion of steel is relatively severe in all the soils. The corrosion of zinc relative to that of steel is considerably greater in the laboratory than in the field. In fact, the laboratory data indicate that zinc is corroded even more rapidly than steel. In the field, the steel corroded more rapidly. Evidently the separation of the electrodes in the cell accelerated the corrosion of the zinc. In the field, the concentration of the alkali at the cathodic areas was greater, so that the zinc hydroxide which precipitated as an adherent film tended to prevent the extension of the anodic areas and the consequent increase in the corrosion current. In the corrosion cell, the electrodes were far enough apart so that the anodic areas would be unaffected by the spreading of alkali at the cathode. The laboratory data might better be taken to represent the behavior of zinc as the anode in a galvanic cell, for example, in the protection of steel cathodically. It is in connection with the protection of steel against corrosion, either as a metallic coating or as a separate anode in connection with cathodic protection, that the behavior of zinc in soils is of the most practical importance.

The relative corrodibility of copper and 60-40 brass in general agrees qualitatively in the two series of tests. However, the marked differences occasionally observed in the corrosion rates of these materials in the field tests, as in soils 23 and 57, are not reflected in the laboratory data. Evidently the electrodes in the corrosion cell are not as much affected by dezincification as in the field tests. It is note-worthy that the superiority of the low-copper brass in the Tidal marsh soil (43), which contains

sulfides, is indicated by both tests.

The corrosion rates of lead at certain of the test sites are subject to the limitation of having been calculated from exposures of only 2 years. If allowance is made for the probable decrease in the corrosion rate with time in these soils, both field and laboratory data may be taken to indicate low rates of corrosion of lead in soils containing high concentrations of chlorides, sulfates, or carbonates.

#### X. SOIL SURVEYS AND TESTS

#### 1. ESTIMATION OF THE CORROSION TO BE EXPECTED IN SOILS

One of the most important considerations in connection with the laying of a pipe line is the amount of protection that should be provided against corrosion. It has already been shown that the various soils differ with respect to their corrosive action on buried metal. Obviously, therefore, an early step should be a survey to estimate the amount of corrosion to be expected in various parts of the territory in which the line is to be laid. If it were true that soils attack metals in much the same way as chemicals do and some one measurable property of the soil could be taken as a criterion of its corrosivity, such a survey would be a comparatively simple matter. Unfortunately, however, the inherent chemical nature of the soil is not always the most important factor involved, and several others must be considered. Furthermore, as soils are seldom homogeneous throughout more than a relatively small volume, a given sample may be more or less corrosive than a similar sample taken only a few feet away. Also, a single test on a composite sample would present merely the average of its components, whereas it is the

maximum corrosiveness of a soil that usually determines its destructive effect. For these reasons, and realizing also that some of the characteristics of the soil are altered in the sampling or testing process, it is evident that the results of a single test may be misleading, and even many tests cannot be expected definitely to indicate the time when the first failure of a pipe will occur. On the other hand, a series of tests on an adequate number of samples properly taken, or a soil survey in which observations are made at frequent intervals, will furnish data from which reliable estimates can be made concerning the performance of a pipe line exposed to the soils thus examined. A soil survey, including the taking of the samples, should preferably be made by an experienced engineer, who should also be responsible for the interpretation of the results because proper conclusions depend not only upon the numerical data, but also upon observations of local conditions, which are often difficult to express definitely. An experienced engineer familiar with soil conditions and their relation to pipe line corrosion may be able to make a better estimate of the corrosive areas from visual observation alone than can be made by a technician skilled only in laboratory methods. The best results are to be obtained by a combination of tests and experience.

## 2. FIELD TESTS (a) RESISTIVITY

The most common criterion of the corrosivity of a given soil is its resistivity. This depends quite largely upon the nature and amount of dissolved salts in the soil, and is also affected by the temperature and moisture content of the soil. Obviously, the resistance of the electrolyte is one of the factors that affect the flow of the current associated with corrosion. If other factors are constant, there is a relation between soil resistivity and corrosion, as indicated in table 89, which presents data obtained by Denison and Ewing [40].

Table 89.—Repairs to pipe lines in different types of soils as related to their resistivity.

[Total acidity 15 to 18 mg =eq per 100 g of soil]

Soil type	Resistivity	Pipe line repaired
	Ohm-cm	Percent
Lordstown fine sandy loam	11,450	3.3
Wooster loam	8,002	6.0
Volusia silt loam	5,473	13.6
Mahoning silt loam	4,903	20.9
Miami silt loam	3,982	22.8
Nappanee clay loam	1,009	57.0

must compensate for inductance and capacity, which vary with frequency. Also, corrections for the temperature and moisture content of the electrolyte must be made. However, as two samples of soil are seldom

identical, and as the factors other than resistivity are rarely constant, an approximate value of resistivity is usually sufficient. The following methods of determining in the field the resistivity of a sample of soil under the conditions existing at the time of the measurement are sufficiently accurate for most corrosion engineering purposes.

(1) Shepard's Soil-Resistivity Apparatus.—Of the several pieces of apparatus used for rough determinations of soil resistivity by corrosion engineers, Shepard's [97] apparatus, as shown in figure 49, is probably most widely used. The apparatus consists of small electrodes attached to insulating rods. The electrodes are connected to a battery and a milliammeter and thrust into the soil to be tested. Shepard describes the apparatus as follows: "The anode is an iron cone of one-half inch (1.27 cm) base and one-quarter inch (6.3 mm) height. The cathode, also of iron, is a frustum of a cone 25% inches (6.7 cm) long, three-quarters inch (1.9 cm) lower base, one-eighth inch (3 mm) upper base. The point is ground or turned to a ½6-inch (1.5 mm) radius. A three-volt battery of two flashlight dry cells is used, the positive pole being connected through a switch to the smaller electrode or anode."

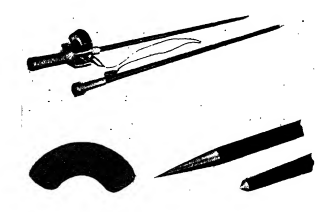


FIGURE 49.—Shepard soil resistivity meter.

A milliammeter having ranges of 25 and of 100 milliamperes is employed. The 25-milliampere range is good for resistivities down to 370 ohm-cm, whereas the 100-milliampere range is good for resistivities between 400 and 100 ohm-cm. The scale of the meter can be graduated directly in ohm-centimeters. A multiplier coil is included in the meter so that the voltage of the battery can be checked from time to time. With such an instrument, the soil resistivity in ohm-centimeters is equal numerically to approximately three times the measured resistance between the electrodes when they are separated in the earth about 8 inches (20 cm) or more. The resistivity meter measures the resistivities of sodium chloride solutions ranging from 100 to 10,000 ohm-centimeters, with an accuracy of about 6 percent.

The accuracy of the reading depends on the condition of the battery and on the resistance of the contacts between the electrodes and the soil.

The apparatus cannot be used in very dry or rocky soil. Because the apparatus measures the resistivity of only a small volume of soil, a single reading may not be indicative of the resistivity of the average soil throughout the region of the test. This objection can be overcome by making several tests in different holes. The apparatus is easily portable and inexpensive, and a large number of observations can be made in a few hours either in the walls and bottoms of trenches or in holes driven in the ground. The apparatus may be purchased from either of two manufacturers or be constructed from directions published by Shepard [97].

(2) Wenner's Four-Terminal Method.—The average resistivity of a large volume of earth can be determined from the surface of the ground by a method developed by Wenner [98] and applied by means of (1) the McCollum earth current meter, (2) the Megger, or (3) by a voltmeter, ammeter, and battery. Four contact points are placed in the earth at equal distances a in a straight line. An alternating or periodically reversed current is caused to flow between the outer electrodes, and the resulting differences of potential between the inner electrodes is observed. If the depth to which the electrodes is inserted in the ground is small compared with the distance a, the effective resistance between the potential electrodes is given by the equation  $R=r/2\pi a$ , where r is the resistivity of a unit volume of soil in terms of the unit in which a is measured. According to Wenner, the measured resistivity depends mainly on the resistivity of the portion of earth situated between the inner electrodes and having a cross section equal to the square of half the distance between the outer electrodes.

According to Rooney [99], who describes apparatus for measuring earth resistivity, the resistivity determined by this method is a general average value in which the resistivity near the line of the terminals is more heavily weighted. He says that, roughly speaking, the effect of material, at a distance (a) equal to the distance between adjacent terminals, is found to be so small that the effect of materials beyond it is negligible.

Biddle [100], who has designed an apparatus for field measurements of soil resistance by the four-terminal method, says that under conditions specified above the resistance measured is that of a half cylinder of length a, and radius 2a. The four-terminal method has been used for studying the changes in soil resistivity with depth and, hence, for determining the desirable length of vertical anodes or the depth at which they should be placed. One corrosion engineer [101], employing this method, uses a B battery for the source of current and reduces polarization errors by using a vacuum-tube voltmeter for the potential measurement.

(3) Electrolytic-Bridge Method.—The electrolytic-bridge method for measuring soil resistivity is employed by the U. S. Department of Agriculture [102]. Fifty milliliters of soil is saturated with distilled water and put into a specially designed vessel containing two electrodes, and the resistance between the electrodes is measured by means of a Wheatstone bridge excited by a vibrator. A telephone receiver is used as a null indicator. By means of tables, the resistivity and salt content are determined from the measured value of resistance. The apparatus is easily portable and can be purchased from an electrical supply house. The determination of the resistivity of the soil when saturated has the advantage of reproducibility of condition and avoids the problem of the proper compactness of the sample. The question as to whether or

not the sample is representative of local conditions remains. Also, if desert soils are involved, the resistivity thus observed may be quite different from that of the dry soil from which the sample was taken.

(4) Radio Balance.—Another method of making a soil-resistivity survey, probably the least accurate of those discussed herein, has been described by Huddleston [103]. This consists in the use of the radio balance, an instrument originally intended for use in locating pipe lines or other buried metal. It consists of a six-tube, portable, impedencecoupled radio receiver and a two-tube oscillator. Both of these instruments are housed in wooden cases with built-in loops and batteries, and are mounted at the ends of the wooden rods, between which the operator walks. The apparatus operates on the principle that conducting materials act as a shield to the propagation of high-frequency waves. Consequently, the presence of conducting materials can be detected by properly designed apparatus. The instrument must be calibrated while holding it in succession over a number of soils of known resistivity. The deflection of the receiving set is proportional to the conductivity of the soil to a depth of about 4 feet. The test requires no excavations and gives a continuous reading. The apparatus is especially useful in approximately locating the boundaries of low-resistance soils, which are usually corrosive. Huddleston reports that 10 miles of right-of-way can be covered in a day by this method. The instrument weighs about 30 pounds.

#### (b) COLUMBIA ROD

The measurements discussed so far indicate only the resistivity of the soil. The currents that accompany corrosion depend not only on the resistivity of the soil but also on the potential of the metal with respect to the adjacent electrolyte and on the limitation of the corrosion currents by polarization films and corrosion products. Several attempts have been made to design soil-testing apparatus that would take account of one or more of these factors. One of them is the Columbia rod developed by Legg [104].

developed by Legg [104].

Figure 50 shows one form of the Columbia rod. It consists of a 1½-inch insulated rod tipped with a cone of steel, a few inches from

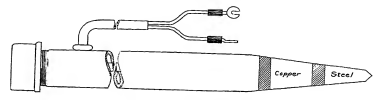


FIGURE 50 .- Columbia rod.

which is an insulated ring of some more noble metal, such as copper. The two electrodes are connected to a sensitive milliammeter, which reads the current flowing when the two electrodes are in contact with the soil. The instrument is calibrated in terms of readings in soils of known corrosivity. The readings are affected by the quality of the contacts between the electrodes and the soil and is satisfactory only for some soil conditions. The changes in the current between the electrodes are indicative of film formation, and the amount flowing

depends on the effective areas of the electrodes as well as on other and more constant factors.

The apparatus could be made with the electrodes on separate rods, as in the Shepard apparatus, and an external battery could be used. This, however, would mask the galvanic effect and the polarization would depend on the applied voltage. The Columbia rod has not been fully developed and probably could be improved. It is not used extensively. Some tests indicate that the readings of the rod are affected chiefly by the resistance of the soil. The amount of soil involved in a single test is less than in a single test by the Shepard apparatus.

#### (c) REDOX APPARATUS

Soil corrosion in the eastern part of the United States, and perhaps elsewhere, is partly due to the depolarizing effects of anaerobic bacteria. The relation of these bacteria to corrosion is discussed elsewhere in this Circular. Starkey and Wight [105] have recently devised a means of determining whether or not a soil condition is favorable to the development of certain of these bacteria. The apparatus consists of two long insulated cylinders, one containing a glass electrode and a calomel electrode, by means of which the pH of the soil solution in a hole in the soil can be determined, and the other containing a platinum electrode and a calomel electrode for measuring the oxidation-reduction potential of the solution. From these measurements it is possible to determine whether or not the aeration and pH of the soil are favorable for the growth of some kinds of sulfate-reducing bacteria. A survey by means of the apparatus would probably indicate the corrosiveness of wet soils, such as swamps, marshes, and first bottoms of streams, but would not identify corrosive, well-aerated soils, except insofar as the corrosivity is indicated by the pH of the soil.

#### (d) IDENTIFICATION OF SOIL TYPES

The U. S. Department of Agriculture has examined and mapped by counties or similar subdivisions about two-thirds of the arable land of the United States, describing the characteristics of each soil horizon, usually to a depth of 6 feet. Each soil that differs chemically or physically from another is given a different name. Insofar as corrosion is dependent on soil characteristics, the name of the soil should indicate its corrosiveness if all the characteristics of the soil are known. However, the corrosiveness of only a few soil types or series has been investigated, and the corrosiveness of a soil series must usually be inferred from its chemical and physical characteristics in the absence of any experimental data. It may be possible, however, to recognize soil types described in Department of Agriculture publications and by associating unknown soils with known soils having similar descriptions, to form some idea as to their corrosiveness. Moreover, as in many regions the same soil types are recurrent, a soil survey along a right-of-way may be helpful in subdividing the territory with respect to its corrosivity. This is illustrated by the work of Denison and Ewing [40], who made a study of the repair records for 200 miles of pipe line right-of-way in Northern Ohio, which contained five parallel lines ranging in age from 26 to 44 years. Some of the physical characteristics of the soils traversed are shown in figure 51. In this study the number of repairs were taken as the measure of the corrosiveness of the soil. It will be shown later that some of the lines were repaired at the same place more than once. The repaired sections were usually given a protective coating

which presumably prolonged the serviceability of the pipe, and thus is equivalent to reducing the corrosivity of the soil. The length of the line repaired depended to some extent on the judgment of the operator. All these things interfere with correlating soil types with corrosion, as expressed in terms of reconditioning.

COLOR	LIG BROWN	7	BRO	WN GRA	YIS	H E	ROW	N GRA		TTLE	2			BRO	WN	
PARENT MATERIAL	, a GI	ACI	AL	TIL	MO'	TTLI	D	\$ 0.00 0.00 0.00 0.00			REAM	GRAY	LACK			Ē
TEXTURE OF SUBSOIL	LIGHT	VERY HEAVY	НЕАVY	LIGHT	VERY HEAVY	неалу	LIGHT	VERY HEAVY OR HEAVY						SANDSTONE	SHALE	
SOIL SERIES	WOOSTER	ELLSWORTH	RITTMAN	CANFIELD	MAHONING	MEDINA	VOLUSIA	TRUMBULL	BRACEVILLE	CHENANGO	CHAGRIN	ноггу	MUCK & PEAT	LORDSTOWN	ALLIS	

FIGURE 51.—Characteristics of soils in the regions of glacial sandstones and shales.

In table 90, in addition to the data on soil types and pipe repairs given therein, are reported the acidity and resistivity of the soil types

so as to give some idea as to their homogeneity.

The significance of acidity and resistivity as measures of soil corrosivity is discussed in another section. It is apparent from the magnitude of the standard deviations of the mean and from the number of samples, that the average values given for the different soils are not equally reliable. Thus the average acidity and resistivity of an extensive soil type, such as Mahoning silt loam, was determined with a relatively high degree of precision because a large number of samples of such soils were taken. On the other hand, less reliance can be placed upon the values for soils present only to a limited extent along the pipe line as they are represented by so few samples.

It will be noted that for three soils, namely, the Clyde clay loam, the Brookston clay loam, and the Crosby silt loam, the data have been separated into two groups, for which, however, the respective acidities and resistivities are practically the same. This separation was suggested by the fact that the extent of repairs differs greatly in areas of each of these soil types, depending upon their position with respect to a certain division point on the pipe line. West of this division point, where the soils are more corrosive, the pipe lines closely parallel a railroad for many miles. It is possible that the accumulation of cinders on the surface of the soils might account for the greater corrosiveness in this region.

TABLE 90, -Summary of data on soils and pipe-line repairs.

	A DOLLS FOR THE TANK HALL OF CHILL OF CHILL FOR THE LY PRESENT	a framer h			The same	Lane				
	Soil	Vumbon	Acidity (r	Acidity (milliequiv- alent/100 g)	Resist (ohm-cm	Resistivity (ohm-cm at 60° F)	Length	Average	gh standard	
Symbol	Symbol Type	of samples	Mean	Standard deviation of mean	Mean	Standard deviation of mean	(in hundred)	of pipe repaired	of percent- age of pipe repaired	of soil bodies
-	2	m	4	70	9	7	œ	6	97	11
A ₁ B ₂ B ₆ 1	Clyde clay loam, first section Clyde clay loam, second section Clyde clay loam, second section Brookiston clay loam, first section Brookston clay loam, second section	23.2 23.2	14.6 14.0 15.0 14.2	1.5 1.9 0.9 2.7	2,758 2,983 9,410 3,051 2,965	224 112 215 600	2,180 505 164 3,231 3,101	46 2 7.0 23 3 34 1 18.4	2022 2022 2023 2023 2033 2033 2033 2033	30 4 1 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2
ට්ට්ට්ට්ට් ට්ට්ට්ට්ට්	Cancerdon siti boam and fine sandy Joan. Canfield siti, Joan. Chaptin siti toam and fine sandy Joan Cindens. Crosby sit I Joan, first section.	8 15 16 19	12.7 18.7 12.6 14.6 22.0	2.1 1.0 1.2 1.2	4,329 9,773 6,184 2,081 4,337	837 1,087 675 160 260	1,324 1,891 907 562 1,659	13.3 6.2 31.7 50 6	11.0 22.5 16.0 16.9	19 20 7 34
SOMOH SOMOH	Crosity silt loam, second loam. Clenango silt loam Ellsworth silt loam Genese silt loam. Holly elay loam.	22566	19.0 9.5 20.4 7.3 26.7	2.5 1.0 2.4	5,660 8,320 7,134 2,661 3,100	1,880 512 763	330 317 655 367 928	3 5 19.6 16.1 33 9 27.7	28.33 14.7 15.0	46 113 171
EEUKK	Lorain fine sandy loam Lorain elay loam Lorains elay loam Milischa en sandy loam Lorae and other shallow limestone soils Lucas fine and very fine sandy loams.	12 23 H	7.0 14.0 16.5	3.2	4,400 3,480 11,450 7,210	321	160 745 290 680 184	28.6 7.1 3.3 11.1 0.0	37.4 5.9 10.5 2.7	420000
M Mu Nu Nf	Mahoning silt loam. Minni clay loam, silt loam, and fine sandy loam. Nutok. Nappanee clay loam. Newton fine sandy loam.	37 17 13 13	18.1 16.8 36+ 17.5 8.0	9.7 3.6	4,903 3,982 2,070 1,009 2,820	475 452 350	5,637 1,671 134 535	20 22 22 23 20 24 25 25 26 26 26 26 26 26 26 26 26 26 26 26 26	13.3 30.6 16.2 10.7	714 mare
A H I I N	Plainfield fine sand includes gravelly phase. Undifferentiated sands and saudy loams. Trumbul leafy foun and silt loam. Till (some Alia and Chenango). Volusia silt loam.	20 e x = 13	6.7 8.0 21.1 9.3 17.2	2.3 5.6 1.0	6,720 2,990 4,455 2,525 5,473	720 920 383	1,160 3,555 610 610 4,105	20.4-3 20.0 13.6-3 13.6	74881 19009	80218
VI WI W	Volusia Ioan Wooster fine sandy Ioan Wooster joan Wowster joan Wauseon fine sandy Joan Wauseon fine sandy Joan "Wauseon like" soils	re 25 23 11	10.4 14.7 15.5 7.5 8.0	1.3	6,023 8,002 1,248 691 691	2,718 1,010	405 278 3,715 240 640	7.1 12.4 6.0 6.3 61.0	23.27	7 4 S - 10

A summary of the data on soils and pipe line repairs is given in table 90. The repairs which have been made in the different soils are expressed as percentage of length per soil body. According to this method of expressing the data, the percentage of repairs (column 9) in the various bodies of a given soil type are simply averaged without regard to the size of the body. The standard deviation of the percentage of pipe repaired in each soil type (column 10) was obtained after weighting the percentages for each separate body according to the

length of line in that body.

The data in table 90 show that, although there is considerable spread in the amount of pipe replaced in various areas of the same soil type, there are consistent differences in the corrosiveness of the various types. Thus among the most corrosive soils must be listed the Clyde clay loam first section (A₁), the soil overlaid by cinders (Ci), the Nappanee clay loam (N), and the "Wauseon-like" soils (Wa). On the other hand, the Canfield (Ca), Lordstown (L), and Wooster (Wl) loams, and the large group of undifferentiated sands and sandy loams (S) must certainly be considered as practically noncorrosive. The certainty with which the corrosiveness of the different soils is known depends upon the extent to which the soils occur along the pipe lines. For instance, relatively little weight can be given to the value for the Lorain fine sandy loam (L), since only one-half mile of this soil was mapped.

Table 90 also shows that the percentage of repairs in small bodies of a given soil type is more likely to differ from the average of that type than are the repairs in larger bodies. Errors in mapping the type boundaries and in locating the places where repairs were made in small bodies of soil introduce larger errors than in a single body of equal area.

The wide spread in the data summarized in table 90 may be illustrated by the repairs in a single soil type. In table 91 are shown the repairs in five bodies of the Lordstown sandy loam. It is easily seen that the repair of 570 feet in the 1,000-foot length of pipe line is inconsistent with the fact that only 380 feet of pipe were repaired in the remaining length of 28,000 feet. In this body where the pipe was repaired, which extended for 200 feet (5 lines), it would appear either that the soil is not really Lordstown sandy loam or the pipe did not require reconditioning solely on account of the corrosivity of the soil. Electrolysis or long line currents may have accelerated the corrosion.

Total length of line	Length repaired	Length repaired
ft 2,000 1,000 10,500 9,500 6,000	ft 0 570 0 0 380	% 0 57 0 0 6.33
Total29,000	950	3.27

Table 91.—Repair data for Lordstown sandy loam.

The distribution of corroded areas as related to soil type is shown graphically in figure 52 for a typical section of the lines. Pertinent data such as the total and repaired lengths of line in each area and acidity and resistivity data are also shown. The marked differences in the corrosiveness of several soil types is immediately apparent. In the areas occupied by the Chenango silt loam (D) and the Holly clay loam (H),

it is seen that as many as three separate repairs have been made on short lengths of the lines. In contrast to these severely corrosive areas are intervening areas of Wooster loam (W1) in which the repairs have been negligible. Similarly, it is seen that the area occupied by the Mahoning silt loam (M) is corrosive, but that few if any repairs have been made in the Canfield (Ca) and Ellsworth (E) silt loams and in the Volusia loam (V1).

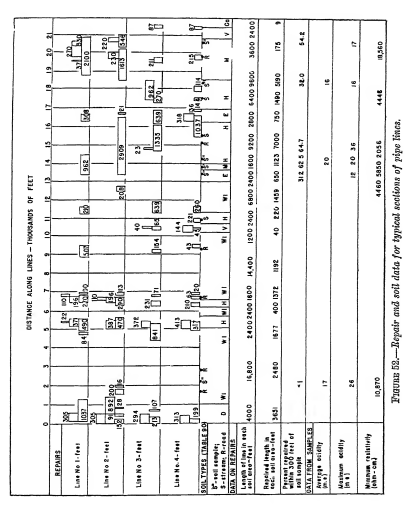


Figure 52 further illustrates the errors, previously referred to, which may occur at the boundaries of the different soils. Thus it is seen that practically all the repairs that have been necessary in the area of

Wooster loam between 1,000 and 5,000 feet have been made at the

boundaries of the adjacent soil types.

In considering further the relation between the various soils and their corrosiveness, it is of interest to compare the repairs made in certain soils with the degree of development of the horizons within the soil profile. In table 92, the typical upland soils of northeastern Ohio, which have developed from sandstone and shale, have been grouped in four vertical columns according to the degree of development shown by their profiles. Within each vertical column the soils are arranged according to the texture of the B-horizon. In table 93 a similar arrangement is shown for the glacial soils of northwestern Ohio, which have been derived from limestone.

Table 92.—Relation between the degree of development of the glacial soils of northeastern Ohio and their corrosiveness.

	Degree of development								
	1		2	;	3	3	Brown No mottling		
Color of surface soil  Mottling of A ₁ horizon	Gray Mottle	ed.	Gray l		Light No mo				
Mottling of B horizon	do		do		Mottled		do		
Mottling of C horizon	do		do		do		do		
	Series	Repairs	Series	Repairs	Series	Repairs	Series	Repairs	
Light B horizon  Heavy B horizon	Trum-	Percent 20.0	Volusia	Percent 13.6	Canfield	Percent 6.2	Wooster	Percent 6.0	
Very heavy B horizon	bull		Mahon- ing	20.9	Ells- worth	16.1			

Table 93.—Relation between the degree of development of the glacial soils of northwestern Ohio and their corrosiveness.

	Degree of development							
	1		2	2	3			
Color of surface soil	Grayish bl	ack	Dark gray		Gray brown			
Color of subsoil	Mottled bl	uish gray	Mottled h	luish gray wish brown	Mottled yellowish brown and yellowish gray			
	Series	Repairs	Series	Repairs	Series	Repairs		
Light B horizon		Percent		Percent	Miami	Percent 22.8		
Heavy B horizon	Clyde	46.2	Brookston	34.1	Crosby	30.8		
Very heavy B horizon		· · · · · · · · · · · · · · · · · · ·		• • • • • • • • • • • • • • • • • • • •	Nappanee	57.0		

It is seen from the tables that the corrosiveness of the soils is related to their stage of development, the least developed soils being invariably the most corrosive. Thus the Trumbull soils, which are mottled throughout the profile and show very little differentiation into horizons, are seen to be corrosive, whereas the Wooster soils, which are well developed, are noncorrosive. It will also be observed that within the vertical col-

umns corrosiveness increases as the subsoil becomes heavier in texture. The relationship between the stage of development and texture of the soils and their corrosiveness can be largely explained on the basis of their average acidity and resistivity. Because of the slight weathering that has occurred in the case of the poorly developed soils, such as those of the Trumbull series, there has been but little tendency for soluble materials to be removed, with the result that the average resistivity of these soils is relatively low. Similarly, the very heavy texture of poorly drained soils accounts largely for their high acidity, the acidity of a soil being a function of its content of colloidal material. Conversely, the high stage of development of the noncorrosive Wooster soils has resulted from the thorough removal of soluble salts, as indicated by their high average resistivity. Aside from the effects of acidity and resistivity, however, it is highly probable that those differences in the physical characteristics of the soils that determine their drainage and aeration have an important bearing on the observed relations.

The correlation of corrosion with soil types is helpful because the U. S. Department of Agriculture has mapped half of the soils of the United States, classified them as to soil type, and described each type. It might be possible, however, to make a simpler correlation by the use of soil series instead of soil types, as usually the different types in a soil series differ only in the texture of the uppermost layer, or A horizon. That is the subsoils are usually the same for most of the soils belonging to the same soil series. It is, therefore, of interest to examine the consistency of corrosion within the series, as it would be desirable, if possible, to take a soil series rather than a single soil type as a criterion for corrosion. To obtain data on this question, specimens of open-hearth iron, wrought iron, steel, and cast iron were buried in two or more soil types belonging to the same series but separated as to location. Unfortunately, some of the alkali soils that were selected were afterward found to differ in salt content. Table 94 shows the results of the tests. As the duration of the exposure in different soil types belonging to the same series differed somewhat, the corrosion losses and pit depths have been reduced to rates per square foot. The table is satisfactory for comparing materials in the same soil series, but should not be used for predicting corrosion for longer periods of exposure.

If allowance is made for the probable spread of data assignable to uncontrolled factors, it will be seen that corrosion within the soil series investigated is reasonably constant. An exception is the Norfolk series where the corrosion in Norfolk sandy loam is different from that in Norfolk sand. The subsoils of these two types are quite different. This serves as a warning that before assuming that corrosion throughout a soil series is the same, one should determine the sameness of the subsoils.

TABLA 94.--Relation of soil series to rates of loss of weight and to weighted rates of maximum penetration of ferrous specimens. Average of 2 specimens, except for the pit cast (A and 1.), the data for which cover 1 specimen only.]

7.9 7.6 7.2 7.8 11.8 8.9 16.9 19.5 18.5 18.3 39.6 13.3 19.0 25.6 19.3 39.5 Aver-age for six speci-mens 39 Weighted rates of maximum penetration in mils per year 12.85 centri-fugal cast iron, 14.8 22.4 19.237.4 41.9 cast 19. 8 œ. 8 25 9 20. 46, North-စ်လက် နှေသပ 25.1 ∞ 29.9 9 0 21.7 43.4 ern east iron, Н 12 82 7.6.18.98 ଞ୍ଚ 36. ଛ 21. 88 6 N 7.3 8.8 10.1 7.2 17.9 26.6 45.4 50.1 12.3 28.3 22.9 28.7 26.1 54.7 cant iron, 5 6 8 €. ⋖ 37.5+ Вевве-10.9 16.0 12.7 13.2 œ mer steel, 80 10.0 7.1 z 19 12 88 37. puddled wrought iron, 26.5+ Hand-А 13.6 7.6 7.5 7.5 9.0 7.5 9.0 12.5 18.0 27.9 9.8 13.9 11.1 6.1 10.4 C 9.7 'n, 20 a17.1+ 38.4+ Open-hearth iron, 14.6 910408 9.9 12.8 14.7 13.7 19.1 ₹ 5 r; ro; 4; ∞ ∞; o; 8 38. 1.86 1.96 2.432.08 Average for six speci-mens 1.05 2.27 2.33 75.88.58.84 0.473.663.943.67 Rates of loss in ounces, per square foot per year Monocast centri-fugal cast iron, 0.45 2.26-57.5 2.49 4.88 2.75 0.50 8883 2.082.742.36 0.87 PD 0.45 ern cast ron, 1.18 2.76 2.8434.88.4.8.4. 1.86 1.84 2.57 2.095.28 5.05 4.57 0.58 .335 .277 .88 South-0.44 3.25 1.10 2.77 3.40 1.942.73 4.13 6.33 3.00 cont ron, ö А ¥ Berse- 5 88.45.83 84.83 mer steel, 0.99 1.90 1.921.60 0.53 1.22 2.192.181.86 3.17 8 88 z e. က် Hand- I puddled wrought а 1.742.30 1.68 0.491.71 2.121.70 2.85 3.68iron, 1.01 27 'n, er. 0.61 .35 .45 .46 .46 Open-hearth iron, 1.13 1.98  $^{2.05}$ 1.7 0.48 1.261.98 1.75 3.34 4.34 2.01 84 ¥ 60 ears 9.27 9.27 9.27 9.24 9.24 8.90 5.93 5.93 fign for test Atlanta, Ga.
Macon, Ga.
Salisbury, N. C.
Raleigh, N. C. Grand Junction, Colo. ....do.... Charlotte, N. C. ....do.... Location Kernell, Calif. Fresno, Calif. Niland, Calif. ...do ...do Billings silt loam (moderate alkali). Billings silt loam. Average..... ....op.... Cecil fine sandy loam. Cecil gravelly loam. Average..... (moderate alkali),
Presno fine sandy loam. (high alkali), Fresno fine sandy loam. Fresno fine sandy loam. Billings silt loam..... (high alkali). Average..... Imperial clay. (high alkali). Imperial clay (moderate alkali). Soil type (low alkali), Noil 101 102 103 104 104 104 104 104 104 110 60 Ξ

7.1	6.5	18.6 18.3 16.4	17.8	86.000 86.000	4.1	0.89.6 6.34.6	8.7	9.6	11.8	4.3	4.3
		- 6 11 12	· Ŧ	× = = = = = = = = = = = = = = = = = = =			:	:	. :		:
13 1	. E	525 - 25 - 25	: <del>.</del> .	=-21- X 12 23 13	: =	852av 0+au	- - -	# :	` :	x rc	72,7 112
9 6		288.57 7.79	36.0	5124-12 5125-12	13	\$  \( \times \) \(	2	F 9	19.7	17.2	7.0
70.70 	5.1	13.1 9.1 14.x	12.1	6.2 1.7 3.0	3.3	6.6.5	65	. ×	9.1	21	72
4.8	4.5	13.1 17.2 13.2	14.6	5.7 2.3 2.0 3.1	3.3	6.7 4.9 6.7 5.7	0.0	10 4 9.6	10.0	3.0	2.7
4.6	5.7	14.7 12.8 19.1	15.5	8,2,2,8	4.1	10.3 8.6 5.5	9.9	9.0 8.0	8.5	2.5	3.3
0.67	0.47	2.98 3.05 2.16	2.73	0.66 .12 .082 .24	0.28	1.32 0.87 .67	0.89	0.45	0.91	0.16	0.30
0.17		3.59		0.50		0.85					
0.72 0.20	91.0	24 2 25 8 23	2ĭ ≈	88.95 88.95	1 20	_° ⊼Ճଘ%	0.63	g :			0.17
Si c		272 e	123	0 % 8 % 8 %	0.23	20 20 20 20 20 20 20 20 20 20 20 20 20 2	- E -	2.16	1 27	5 12 M	<u>e</u> ;
79.0	00 0	222	2.15	6.75 5.75 5.75 5.75 5.75 5.75 5.75 5.75	0.33	0 92 17. 17.	0.83	===	0.80	0.16 82	31 C I
0.6 8.	0 10	2 06 2 16 2 11	2.05	5,55%	₩.	. 17.0 17. 18.	0.77	널릴	17	0 28	급 :
5 K	=======================================	2 2 2 2 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3	2 13	6 8252	- F	2328	× c	2.2	0.75	27	31 C
¥8 ==	:	252		===57.		급드급급 _ 요원26 2	:	5 tg 75 dg		22	<u>.</u>
Memphis, Tenn11.455 Vicksburg, Miss		Buttonwillow, Calif. Los Banos, Calif. Tranquility, Calif		Maoon, Ga. 17 Pensacola, Fla. 11 Tampa, Fla. 12 Jacksonville, Fla. 12		Meridian, Miss. Shreveport, La. Troup, Tex.		Los Angeles, Calif Bakersfield, Calif		Milwaukee, Wis Springfield, Ohio	
Memphis silt loamdododo	Average	Merced silt loam	Average	Norfolk sandy loam Norfolk sand do Norfolk fine sand	Average	Susquehanna olaydodoSusquehanna silt loamSusquehanna fine sandy loam.	Average	Hanford fine sandy loam Hanford very fine sandy loam.	Average	Miami clay loam	Average
115		23 116 117		3128		25 123 125 125		132		26	
814786°—48——14											

 $^{\rm a}$  +indicates 1 or more punctures due to corrosion. b D=Destroyed by corrosion. • Data for 1 specimen only. The other was destroyed by corrosion.

## 3. LABORATORY TESTS (a) RESISTIVITY

The conductance of most soil is almost entirely electrolytic. The resistance of a soil, therefore, depends on the moisture in the soil, the salts in solution, the distribution of the moisture, and its temperature. The compactness of the soil affects its resistance, and in a measurement of a small volume of soil containing stones or gravel, the result may be affected by the distribution of this inert material. If direct current is used, the results may be affected by polarization and endosmose. The National Bureau of Standards [106] made a rather extensive study of soil resistivity in the course of its study of stray-current electrolysis, and for several years made soil-resistivity measurements on samples of soil. These, as received from the field, were placed in a glass-lined cylinder having a volume of about 800 milliliters and provided with a movable piston by which the sample was subjected to a pressure of 50 pounds per square inch. Measurements were made with a voltmeter and ammeter, using 60-cycle current. At 50-pound-per-square-inch pressure, the resistivity of most soils was found to be about that of the undisturbed soils.

Although such measurements may have shown the resistivity of the soil at the time the sample was taken, the results were questionable because another sample taken at a different time might have a very different resistivity because of a different moisture content. Later soil-resistivity measurements were made with the soil sample saturated, using a Wheatstone bridge and a 1,000-cycle current. From the resistance, the resistivity at 60°C was calculated. The volume of soil tested was 46 milliliters. From the standpoint of the soil scientist, the change in method had several advantages, although the results were in many cases not representative of the resistivity of the soil in the field at any time, and the chances of the sample being representative decreased with the decrease in the volume of the sample tested. As the resistivity of the soil in the field may vary considerably from point to point and from time to time, and as resistivity is only one of the characteristics which influence its cor-

rosivity, an approximate value is sufficient for most purposes.

Another container used at the Bureau consists of a Bakelite frame with removable iron electrodes at the ends and having a cross section of 5 square centimeters and a length between electrodes of 5 centimeters. The resistivity of soil in this frame is equivalent to its resistance. Another frame was constructed having a cross section of 30 square centimeters and an inside length of 3 centimeters, so that the resistivity of the soil equaled 10 times the observed resistance. These containers were placed on glass plates, and measurements were made with the alternating current bridge. Wahlquist [101], for some of his field measurements, used a wooden box lined with waxed paper. The box had a length of about 8 inches and a cross section of about 4 square inches. Current terminals were placed at the ends of the box, and point potential terminals were so spaced near the center that the resistivity of the soil equaled the resistance, which was measured by means of a battery, milliammeter, and vacuum-tube voltmeter. The separation of the current and potential electrodes avoids the inclusion of polarization effects at the current terminals. However, the voltmeter terminals may differ somewhat in potential. This source of error may be avoided by applying different currents and dividing the change in voltage by the change in current. When the measurement has been completed, the soil is slipped

out of the box and wrapped in the waxed paper for later reference. The effect of compacting the soil in a test box is shown in figure 53.

The effect of the moisture content on the resistivity of a clay soil is shown in figure 54. Other kinds of soils would yield different curves because of their different salt content and moisture equivalent, as is shown in tables 20 and 21. The effect of temperature on the resistance of one soil is shown in figure 55. For temperatures between 0° and 25°C the relation between soil resistance and temperature is given by the equation  $R_{15.5} = R_t(24.5+t)/40$ , in which  $R_{15.5}$  is the resistance at 15.5°C (60°F) and  $R_t$  is the resistance at the temperature t °C.

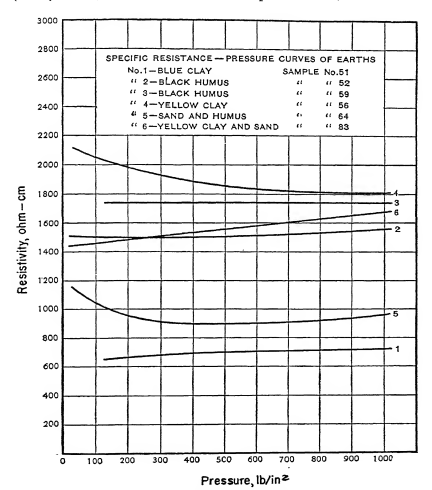


FIGURE 53.—Specific resistivity of soil at different pressures.

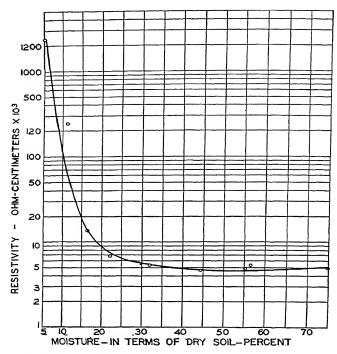


FIGURE 54.—Effect of moisture content on soil resistivity.

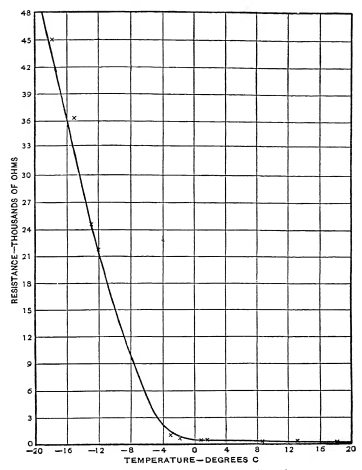


FIGURE 55.—Effect of temperature on earth resistance.

# (b) HYDROGEN-ION CONCENTRATION AND TOTAL ACIDITY

Holler [107], as a result of a study of hydrogen evolution from iron filings in contact with soil in the absence of oxygen, found a logarithmic relation between the total acidity of the soils at the Bureau test sites and the volume of hydrogen evolved and suggested that the corrosiveness of soils in humid regions might be predicted from their acidity. Because of the very slight dissociation of the soil acids, the pH value of a soil may offer no indication of the capacity of the acidic material to prevent the formation of a protective layer of hydroxides that tends to form as a result of initial corrosion. It has been noted that with a definite oxygen concentration the rate of corrosion is determined by the total quantity of ionizable hydrogen that comes into contact with the metal surface, rather than by the hydrogen-ion concentration of the corrosive medium. Unless the quantity of acid that comes into contact with the metal surface is sufficient to prevent the formation of protective hydroxide films, these films will tend to form regardless of the H-ion concentration. In order to determine the effect of soil acidity on corrosion in aerated soil, Denison and Hobbs [39] made up a series of synthetic soils differing chiefly in total acidity. They found that, under the conditions of their test, the rate of corrosion was roughly proportional to the total acidity of their soils. A somewhat less definite relation was found between the rate of pitting and the acidity of soils from Bureau test sites.

The determination of total acidity of soils is somewhat difficult because of the slight solubility of soil acids. The details of the procedure followed in determining acidity by the "difference" method as described by Denison [108] are as follows. The capacity (T) of the soil to absorb exchangeable bases was first determined. A 25-gram sample of soil was placed in a 250-milliliter Erlenmeyer flask with about 1 gram of calcium carbonate added. The soil was treated with 150 to 200 milliliters of 1 N NaCl that had been heated to 80 $^{\circ}$  to 90 $^{\circ}$  C. The mixture was maintained near that temperature for 1 hour, with occasional shaking. The mixture was then poured into a 400-milliliter bottle and shaken overnight in an end-over-end shaker and also for 2-hour periods on 2 succeeding days. After standing another night the extract was decanted and filtered into a 1-liter beaker. Then 200 milliliters of salt solution was added to the soil. The mixture was shaken 1 hour, allowed to settle, and the extract then decanted through the filter as before. This process was repeated until 1 liter of extract was obtained. For the final filtration, the entire quantity of soil was poured on the filter and washed. The extract was then mixed thoroughly, and the calcium in a 400-milliliter portion was determined gravimetrically. From the value thus obtained, the quantity of calcium equivalent to the solubility of calcium carbonate in 1 liter of 1 N NaCl was subtracted.

The quantity (S) of replaceable calcium in the soil was determined in a manner similar to the determination of the exchangeable base capacity, except that no calcium carbonate was added and the shaking was limited to 18 hours. For those soils in which calcium carbonate occurred naturally, a correction was made for the calcium that had been dissolved as carbonate by the salt solution. This quantity was calculated from the amount of bicarbonate in solution, which was estimated by titrating an aliquot portion of the extract with standard acid to the color change of methyl orange. The quantity (T-S) of exchangeable hydrogen or the total acidity of the soil was obtained by

subtracting from the exchange capacity of the soil the absorbed calcium found to be present.

Ewing [109] devised a shorter method that duplicates the results of Denison's method within about 15 milligram equivalents per 100 grams of soil. Ewing's method is as follows: Two 5-gram portions of the airdry soil, previously pulverized and passed through a 20-mesh (per linear inch) sieve, are placed in two 25- by 200-millimeter test tubes, and then 25 milliliters of normal NaCl solution is added to each tube. One milliliter of 0.2 N Na₂CO₃ solution is pipetted into one tube and 2 milliliters into the other. (One milliliter is equivalent to 4 milliequivalents of acid per 100 grams of soil for the 5-g sample.) The pH of the more alkaline solution is then determined. If the pH is found to be below 8, 2 milliliters more of the carbonate solution is added to each tube and the process repeated until the pH of the more alkaline solution is above 8. The tubes are allowed to stand, with occasional shaking, until the solutions are in equilibrium with the soil and show no further changes of pH with time. The pH of the solution in each tube is then determined. Usually the resulting pH values are so near to pH 8 that by interpolation or extrapolation the amount of alkali required to bring the soil to pH 8 can be determined. For example, if 12 milliequivalents (3 ml) of the carbonate solution give a pH of 7.7 when in equilibrium with the soil and 16 milliequivalents (4 ml) give a pH of 8.1, the acidity of the soil will be 15 milliequivalents per 100 grams. The titration curve is assumed to be a straight line through the range of interpolation and extrapolation.

Table 95.—Repairs in soil types as related to total acidity.
[Resistivity 4,000 to 5,000 ohm-centimeters]

Soil type	Total acidity (mg-eq/100 g soil)	Pipe line repaired
Wauseon fine sandy loam Caneada silt loam a Miami silt loam b Mahoning silt loam Trumbull clay loam c Crosby silt loam, first section	12.7 16.8 18.1	Percent 6.3 13.3 22.8 20.9 20.0 30.8

The relation of the total acidity of the soil to its corrosivity is indi-

^a Includes fine sandy loam. ^b Includes clay loam and fine sandy loam. ^c Includes silt loam.

addition of water to the soil.

cated by table 95. Of course, the corrosivity of most soils is influenced by several of their characteristics, some of which are not closely related. Consequently, in many cases there is no close correlation between a single soil characteristic and corrosion. For the northern Ohio soils previously discussed, Denison and Ewing found that the corrosivity could be roughly expressed by the equation P=7500(A-5)/R, in which P is the percentage of pipe repaired, A is the total acidity in mil equivalents of hydrogen per 100 cubic centimeters of soil, and R is the soil resistivity in ohm-centimeters. Although the relation between hydrogen-ion concentration and total acidity that has been discussed affords a sufficient explanation for a lack of correlation between hydrogen-ion concentration and corrosion, several factors that affect the determination of pH values at the National Bureau of Standards and elsewhere may also

have affected the results. One of these was the use of a colorimetric method for pH determinations, which in most cases necessitated the

The difficulty of obtaining a clear soil solution or of matching colors when the solution was not clear sometimes may have resulted in inaccurate determinations. More important, probably, is the fact discovered by Romanoff [47] that if certain soils are air-dried prior to the determination of their hydrogen-ion concentrations, as is the practice of the Department of Agriculture and the National Bureau of Standards, the pH values undergo changes due, probably to the oxidation of sulfides in the soils. The total acidity of the soil may change also. It is possible, therefore, that in some cases the pH values used in the correlation of pH values and corrosion were incorrect. The changes referred to occur in some poorly aerated soils containing active sulfate-reducing bacteria. The hydrogen-ion concentrations of other poorly aerated soils are unaffected by aeration. When conditions permit, the pH values of the soils at the National Bureau of Standards test sites will be redetermined without exposing them to the air, but it seems probable that only a few changes in the published values will be necessary.

# (c) PUTNAM'S DECOMPOSITION POTENTIAL TESTS

A test that has been used extensively in the West is one devised by Putnam. This test has undergone several changes with respect to apparatus and the way in which the data are treated. The original test [110] was one of the earliest attempts to determine soil corrosivity and to associate corrosion with the electrical condition of the corroding metal.

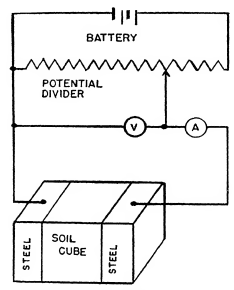


FIGURE 56.—Putnam apparatus for testing for potential corrosivity.

The latest modification [52] takes account of the resistivity and acidity of the soil and of polarization. There are some, however, who doubt the validity of the theory upon which the test is founded.

The test involves the determination of the resistivity and pH of the

soil and the value of current under certain experimental conditions which Putnam calls the "potential corrosivity" of the soil. The circuit for determining potential corrosivity is shown in figure 56. Two polished-steel electrodes having faces 1 inch square are held in a bakelite case (not shown) with their faces 1 inch apart to form cubical cell 1 inch on a side. The soil under test is compacted in this cell under a pressure of 500 pounds per square inch. The resistance of the soil cube is determined by measuring the resistance between the electrodes with an alternating-current Wheatstone bridge. A potential difference of 1.4 volts (causation potential,  $E_c$ ) is then impressed between the steel electrodes by means of a battery and potential divider as shown. After the current has reached a practically steady value, usually after about 5 minutes, its value, C, is read. The equation  $C = (1.4 - \dot{E}d)/R$  gives the value of the current in terms of the causation potential (1.4 volts), the decomposition potential, Ed, and the resistance, R, of the soil cube. The value of C in milliamperes is called the potential corrosivity. Values of pH are determined colorimetrically in a neutral 5-percent potassium chloride solution. By correlating these values with the National Bureau of Standards soil-corrosion data, Putnam derived several empirical relationships for determining such quantities as the loss per square foot, the average maximum pit depths, and the total number of leaks per mile in a given time.

# (d) WILLIAMS-CORFIELD OR NIPPLE-AND-CAN TEST

A more frequently used and simpler test, which, however, has less theoretical background than Putnam's, is the Williams-Corfield [54], or nipple-and-can, test. Its chief claim for consideration is that the corrosion index obtained by its use has been correlated with a large amount of corrosion experience. A 4-inch length of 34-inch sand-blasted iron pipe is carefully weighed, and a rubber stopper is inserted in one end so that it protrudes one-half inch. It is then placed in the center of a pint friction-top tin can, with the stopper resting on the bottom, and the remaining space is filled with a water-saturated sample of the soil to be tested. The pipe is connected to the positive side of a 6-volt storage battery, and the other side of the battery is connected to the can. Several tests are usually run in parallel. Corfield uses a clock to disconnect the battery after 24 hours. The pipe is removed, cleaned, and reweighed. The loss of weight in grams is the corrosion index. Soils are grouped as to corrosivity as shown in table 96.

Corrosion index	Corrosiveness
8 grams and over	Very bad
2 to 2.99 grams	Bad
1 to 1.99 grams	Fair
Less than 1 gram	Good

Table 96.—Corfield corrosivity index.

Figure 57 shows Corfield's apparatus. Corfield tested about 8,000 samples of soil from Los Angeles, Calif. and vicinity by this method to make a soil-corrosivity map of the territory under his supervision. The method has been used extensively by others and appears to be reasonably satisfactory for soils such as occur in California. There are few checks on the reliability of the method when applied to the acid soils of the East.

Among objections to the test are (1) the applied voltage is much greater than those occurring in soil corrosion, and (2) the loss of weight is caused by and is proportional to the current which flows. The current is controlled by the resistance of the soil and by polarization, which in turn is a function of the current. However, Corfield has shown that the same sort of data are obtained when a two-volt battery is used, and the decrease in current due to polarization seems to be a characteristic of the soil. The advantages of the test are ease of manipulation, speed (as many tests can be run in parallel), and the extensive correlation that has been made with field experience.

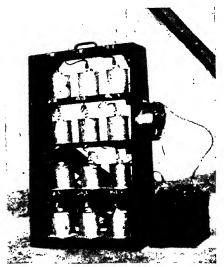


FIGURE 57 .- Corfield apparatus.

### (e) DENISON CELL TEST

Denison's cell test, described on page 181, was devised primarily for studying the general phenomena of corrosion but has been used to a limited extent to determine the relative corrosivities of different soils. The test more nearly simulates actual soil conditions than the tests so far discussed and yields more reproducible results. On the other hand, it requires greater care, more apparatus and more time than the other tests.

### 4. VALUE OF SOIL-CORROSIVITY TESTS

Experienced pipe-line operators can often identify a considerable percentage of the corrosive-soil conditions in a given area, such as swamps, alkali knolls, adobe soils, or alkaline soils without tests. The question is often raised, therefore, as to whether or not the various tests that have been developed for determining the corrosivity of soils are capable of reliably identifying a sufficiently greater percentage of corrosive conditions to warrant their use. Another important question is the relative

value of the different tests. At first thought, it might appear to be a simple matter to answer these questions by comparing the results of laboratory or field tests on individual soils with the amount of corrosion observed on pipes laid in the same soils. As a matter of fact, however, the problem is not so simple. The results of such a comparison will depend to a certain extent upon what criterion is adopted for judging the amount of corrosion. Maximum pit depths, leaks per unit length, number of replacements, or estimated condition of the pipe are the principal criteria that have been used for this purpose. The element of time must also be taken into account because corrosion does not, as a rule, proceed at a uniform rate, and the pipe may not have the same wall thickness in different parts of the line. Also, the relative corrosiveness of various soils is not the same for coated pipe or with respect to their destructive effect on various coatings as it is with respect to bare pipe. In general, it can be said that each of the different tests can identify certain corrosive conditions in the soil but that no single test can identify all places where corrosion is likely to occur. Also it is probable that their relative effectiveness will differ in different localities. Thus, tests of acidity are useless in alkaline soils. Resistivity tests are not very effective if the resistivity is high, and Denison found it advisable to modify his method when testing muck and similar wet soils. These facts should be kept in mind in considering the following comparisons reported in 1939 by Logan and Koenig [111].

In 1937 the Magnolia Pipe Line Co. uncovered approximately 25 miles of 8-inch asphalt-coated steel line in southeastern Texas, measured the depths of the deepest three pits on each joint of pipe, and estimated the condition of each joint. The soils along the right-of-way were mapped according to the methods of the United States Department of Agriculture. 250 soil samples were taken at pipe depth, parts of which were sent to the National Bureau of Standards for test. The resistivity of the soil was determined at 200-feet intervals by means of the Shepard resistivity apparatus. At the National Bureau of Standards, the hydrogenion concentration, total acidity, alternating-current and resistivity at saturation were determined for each sample. The small size of the samples made it impossible to run the Corfield test with full-size apparatus but small-scale apparatus was used that produced the same

current densities as the larger apparatus.

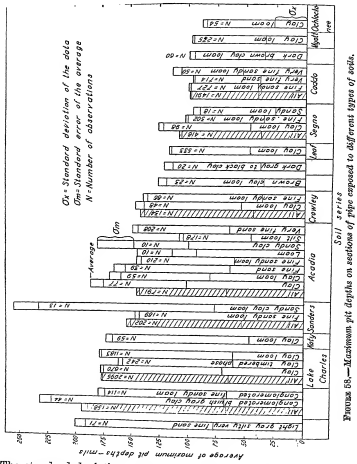
The Columbia rod test was simulated by placing soil in a cell having a copper and a steel electrode. A milliammeter was used to measure the current developed. The Putnam test was simulated insofar as determination of the potential corrosivity, that is, the current furnished by the cell, was concerned, but this potential corrosivity was not reduced to equivalent pit depths by means of Putnam's curves and equation. Therefore, the data do not indicate the reliability of the Putnam test, as described above. The Denison test was also run according to the practice of Denison at that time.

An examination of the data on the pipe indicated that there was a fairly close relationship between the estimated condition of the pipe, the number of punctures, the depth of the deepest three pits, and the depth of the deepest pit on a joint of pipe. For convenience, the depth of the deepest pit was taken as the criterion of corrosion. The data for the various soil corrosivity tests were correlated with the condition of

the pipe as indicated by this criterion.

## (a) CORRELATION OF SOIL TYPES WITH PIT DEPTHS

Figure 58 shows the extent to which the soil series and types were indicative of soil corrosivity. The cross hatched columns represent the averages of all the maximum pit depths in the soil series. The number of observations, the standard deviations of the data, and standard errors of the averages are shown. Most soil series were represented by two or more soil types which are represented by the open columns. Thus the figure shows the variations in the corrosiveness of the soil types within the series.



The standard deviations show how widely the individual pit depths differed. As has been noted, the several soil types of a series usually do not differ greatly in corrosiveness. Figure 58 shows that some soil series are definitely more corrosive than others. The differences shown

are those for soils within 25 miles of each other and do not represent the differences between soils in different parts of the country.

Figure 59 shows the pit depths on sections of pipe adjacent to the spot where the soil was sampled to identify its type. It will be seen that the averages of the pit depths on the pipes in the different soils differ widely and that some soil types are much more corrosive than others. However, the range in pit depths in any one soil type is very great. This should make it evident that the examination of a single joint of pipe in a soil type is likely to give a very misleading idea of the corrosiveness of that type of soil. A large number of observations are necessary for the determination of the corrosiveness of a soil type or the condition of the pipe line in even a single type of soil.

The descriptions of the soil textures near the top of the figure 59 refer to the textures of the soil samples taken at pipe depth. It will be noted that these textures of the soil at pipe depth differ from those given below in the name of the soil type. This is because the latter refer to the textures of the A horizons. The stars indicate the maximum pit depth on the pipe nearest the soil sample for which the texture was determined.

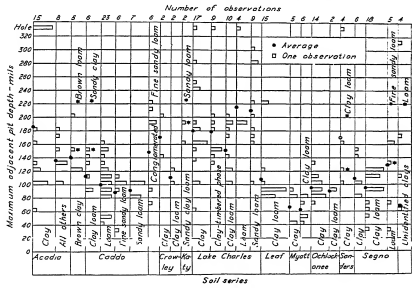
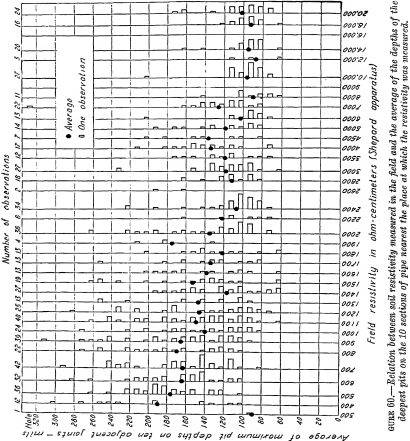


FIGURE 59.—Maximum pit depths on sections of pipe adjacent to places at which soil samples were taken.

## (b) CORRELATION OF CORROSION WITH FIELD RESISTIVITY

(1) Shepard's Soil-Resistivity Measurements.—Measurements of soil resistivity in the field were made at intervals of 200 feet, by means of Shepard's apparatus. These measurements might be taken as representative of soil conditions along ten 20-foot lengths of pipe, and on this assumption the resistivity of the soil should be compared with the average of the maximum pit depths on the 10 sections of pipe nearest the sampling point, as is done in figure 60. The range of pit depths

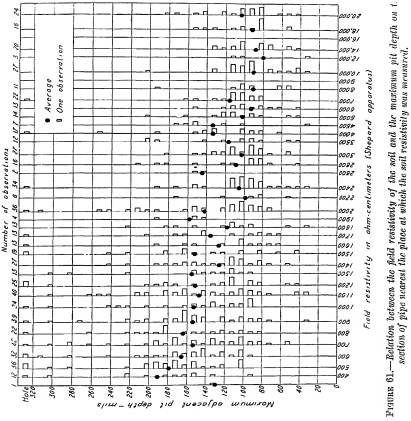
for each value of soil resistivity is so great that it might be thought that the soil sample did not represent soil conditions to which all of the 10 lengths of pipe were exposed. Consequently, a correlation was made between the resistivity of each soil sample and the depth of the deepest pit on the joint of pipe nearest the sample. The result is shown in figure 61. The correlation is no better than that in figure 60. Each figure shows that the average maximum pit depth tends to decrease as



the resistivity of the soil increases, but that there is a wide range of pit depths associated with each value of resistivity. Evidently other factors affect the pit depths.

It might be thought that the poor correlation between pit depths and soil resistivity is obtained because pit depths are not indicative of the corrosivity of the soil. Many pipe line operators divide their used pipe into classes, or conditions, based on the amount of work required to recondition it. One of these classifications is as follows: Condition 1, no pits deeper than 120 mils, no repairs required; condition 2, pit depths ranging between 70 and 260 mils, not more than 25 spot welds required; condition 3, numerous pits but generally not deeper than 40 mils and not more than three punctures; condition 4, junk.

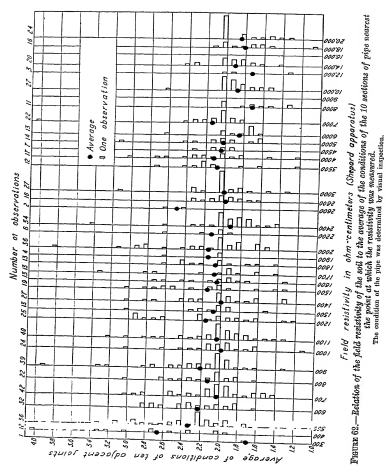
The pipe on which pit depths were measured was classified on this basis and figure 62 constructed. It is very similar to figure 61 and



num pit presents ondition lengths between ordinate istivity. Ind soil dgment

indicates that the two criteria for soil corrosivity, namely, maximum pit depths and pipe conditions, are equally good. Figure 63 presents another way of studying the relation of soil resistivity to the condition of the pipe. The ordinate on the left indicates the number of lengths of pipe in each condition and enables one to study the relation between individual pipe condition and soil resistivity. The right-hand ordinate applies to the average of the pipe conditions (dots) for each resistivity. This curve shows little relationship between pipe condition and soil resistivity, Since the condition of the pipe was based on the judgment

of one or more inspectors and not on measurements, it is not surprising that the assigned condition of the pipe does not accurately represent the corrosivity of the soil.



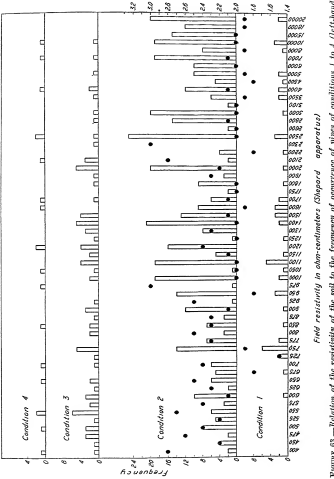
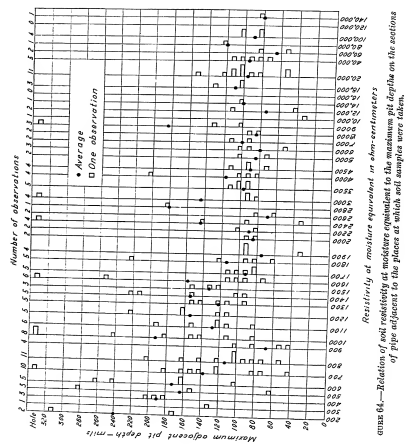


FIGURE 63.—Relation of the resistivity of the soil to the frequency of occurrence of pipes of conditions 1 to 1 (left-hand ordinate) and to the average of the conditions of the pipes (right-hand ordinates).

# (c) CORRELATION OF PIT DEPTHS WITH SOIL RESISTIVITY AT MOISTURE EQUIVALENT

In view of possible differences in soil resistivity due to variation in moisture content, laboratory tests of the soil samples were made at moisture equivalent, which on the average represents the normal moisture content of the soil, to see whether or not this would yield a better correlation. Figure 64 shows that the correlation is, if anything, somewhat poorer, but this may be due to the smaller size and number of samples.



# (d) CORRELATION OF OTHER SOIL TEST DATA WITH PIT DEPTHS

Correlation curves were plotted, using other criteria for corrosiveness, with similar results. Figures 65 and 66 show that the average results for all methods are similar and indicate no close correlation between any of the criteria for soil corrosiveness and pit depths. These graphs indicate

that the results of individual tests are not reliable criteria of soil corrosiveness. The explanation may be that several factors influence the corrosivity of soils and their relative importance differs for different soil conditions.

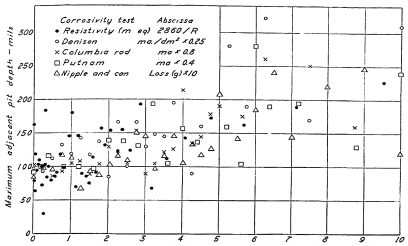


FIGURE 65.—Comparison of corrosivity of soil as indicated by the maximum pit depths on sections of pipe adjacent to places where soil samples were taken with results of various soil tests.

See key at top of illustration for units for the abscissa.

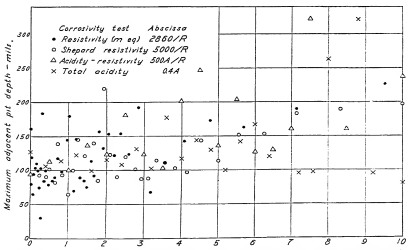


FIGURE 66.—Comparison of corrosivity of soil as indicated by the maximum pit depths on sections of pipe adjacent to places at which soil samples were taken with results of various soil tests.

See key at top of illustration for units for the abscissa.

التانية المالية

For most purposes, it is unnecessary to estimate just how deep the maximum pit at some spot will be at a given time. What is more generally needed is a rough estimate of the average relative corrosiveness of the soils. All that is attempted by many corrosion engineers is to separate the soils or areas traversed by their lines into four or five classes with respect to corrosiveness because it is impractical to provide more than that number of degrees or kinds of protection for their lines. It can be shown that many of the tests under discussion are fairly satisfactory for this purpose.

# (e) RELATIVE MERITS OF TESTS FOR SOIL CORROSIVITY

To determine the relative effectiveness of the tests in correctly grouping soils as to their corrosiveness, the range of pit depths was divided into five groups. The soil series were divided into five groups on two bases: (1) on the basis of the average of the pit depths on the pipe line within each series, and (2) on the basis of the average of the maximum pit depths on the sections of the line adjacent to the places where the soil samples were taken. The range of the results of each soil test method was also broken up into five groups. It should be pointed out here that the choice of all the group boundaries was entirely arbitrary and that it is possible that the relative merits of the tests would appear different if different group boundaries were used. The apparent effectiveness of one or all of the tests might also be improved by a better choice of group boundaries. Nevertheless, the groupings used will give a rough idea of the usefulness of the tests. After the group boundaries were fixed, the soil series were placed in the proper groups as indicated by the pit-depth measurements and by the results of each test and secondary tables were made to show how nearly each soil was placed in each test in its proper place as indicated by pit depths. Table 97 shows four comparisons of the test methods on the basis of the average of the maximum pit depth on the joint of pipe adjacent to the soil sample. Probably because the accuracy of the test results improves with the number of samples tested, the table shows that the percentage of the pipe line correctly assigned as to corrosion is, for most tests, somewhat greater than the percentage of the soil correctly assigned.

Many pipe line operators would be satisfied if the test came within one group of correctly classifying the soils as to corrosiveness. Table 97 shows that all but one of the tests so assigned the soil along at least 86 percent of the pipe line. The test which was least satisfactory on this basis was unsatisfactory with respect to the soil series, which was occupied by 34 percent of the pipe line. None of the tests appears most

satisfactory for determining the corrosivity of all soil series.

Another similar set of comparisons based on the averages of the maximum pit depth on each joint of pipe in a soil series yields somewhat higher percentages of correct assignments for most of the tests. Table 98 shows comparisons of the test methods on two bases for each soil series and the relative values of the tests for all the series without regard to the extent of series or the number of samples tested. It is interesting to note that no one test ranked all the soils correctly or yielded the most accurate ranking for all soils. It is also of interest to note that one soil, Myatt, was apparently not correctly ranked by any of the tests and most of the tests missed the correct ranking very badly. It seems probable that the pitting in this soil was affected by some factor other than soil corrosivity, since all the tests underestimated its corrosivity. It is

TABLE 97.—Comparison of test methods on the dasis of averages of the maximum pit depths on the joint of the specimen adjacent to the soil sample.

Soil series	Percentage of line	Shepard resistivity meter	Resistivity at moisture equivalent	Total acidity	1000 Xacidity Resistivity	Modified Columbia rod test	Putnam test	Nipple-and-	Denison electrolytic cell
		DEPARTU	DEPARTURE FROM CORRECT ASSIGNMENT	ORRECT AS	SIGNMENT				
Aeadis. Brown. Caddo. Congionerste. Crowley.	12.96 0.38 24.43 2.59 2.20	77077	T7°77	07770	07070	1 1 000	77007	00078	70070
Dark brown Lake Charles Lent Light gray Mystf	0.98 34.32 8.73 1.16 4.51	7007	<del>+++</del> +	01540	07477	00001	00008	08000	00008
Ochlockonee Segno	0.88 6.85	°7	++3	++3	H ⁰	00	+10	+10	+10
		PERCEN	PERCENTAGES OF CORRECT ASSIGNMENT	ORRECT AS	SIGNMENT				
Percentage of soils		36	17 26	22.33	42	75 91	58 73	54	67 73
19d	RCENTAGES	WITH AN E	RROR OF NO	T MORE T	PERCENTAGES WITH AN ERROR OF NOT MORE THAN ONE CORROSION GROUP	ROSION GR	tour		
Percentage of soils.		95	67 87	88.88	888	92 97	92 95	22.8	92
								The state of the s	

	arrangement of soils in the sante on and	s and the same, of arach, of corrosterness.
	" basis of correct	
s of testion wile.	Telline Guine C	11.
mparison of method		-
PABLE 98, Co		

n bic	Devia-	mumixam assussed		070	000	707	+ ₂	I 0 ;	= ,
Denison electrolytic cell	Q.	mumizam tasaiba	By	111	+17	0007			o -
Gen		saigned rank		000	8	104	77	6710	:
4	Devia.	mwmizsm szsvs diqeb d	By	1727	09	7000	7	344	
Nipple-and-	De	adjacent maximum t depth	DA BA	1770	+1	4000	7	5 5 7 7 7 7	
ig ig		aigned rank		0007	7	20-98	· ~	₩∞	į
E	Devia- tion	average maximum it depth	B.	7777	1 1	0000	7	20°E	<del>-</del> #
Pulmam test	- Sa	mumixam tasasiba y diqeb ti	B	### ñ	169	7000		70#	73
Puf		dası bəngis	v	8117	-44 0	0.00	G (	. O1 10	-
문흥글	Devia- tion	mumizam egsta y t depth		7777 1177	0 -	700	ç +	178	87
Modified Columbia rod (est	Ğ,	mumixam tagab tic	B	11759	7 7	700	F T	24.	73
	1	saigned rank		00-0	00 r.c	11 6 7	- 61	4 :	i
Resistivity	Devia- tion	mumixam erasiona vi digida depth	1	2222	o 7	7777	9	78	
	2,2	mumixam dassina us Atqəb tiq	I	-11n	7 7	++1+		3.1	7
Re	!	Agar bangiss#	- 1	7:0-1:00	0 01	2000	7	4 :	
Total aridity Devin-	Devin- tion	By average maximum pit depth	:   F	16767	7	1 + 1 +	110	47	∞
	2-	By adjacent maximum pit depth	ī		0	1+1+3	8	74	∞
		Ansı bəngissA	2	4 6 6 9	and 2	50000	=-	- [	
Resistivity at maistare equivalent	Devin- tion	By average maximum git depth	1	4777		,00×	1-6	. 92	, rc
	=	By adjacent maximum pit depth	9-	0113	77	1008		88	9
====	,	Assigned rank	20	11 8 7	4 6	01 10 10	60 00	:	
Eş.	Devia-	By average maximum pit depth	0	4111	121	1000	70		9
Shepard resistivity meter	ă J	By adjacent maximum pit depth	173	7777 111	177	00%	00		2
		Ansı bəngissA	6;	4	C1 00	929 10	cc 10	-	
or order, of corrosi- veness		By average maximum fit depth	97	3and 4 11 8	and 4	252	12	<del>.</del>	
5 g 7		By adjacent maximun pit depth	Ξ×	401-	10 3	200	20.00	<del>:</del>	
		NITH NOTES	AcadiaBrown	Caddo	Dark brown. Lake Charles	Light gray Myatt	Segno	Rank, or order, of effectiveness of	

difficult to see, however, what outside factor would greatly affect the average condition of the pipes at 10 locations or the average corrosivity of 10 soil samples. Inspection of the last two lines of table 98 will bring out the fact that several of the tests are of nearly equal reliability, but it seems probable that the relative values of these tests might be different if soils from some other part of the country had been tested.

Although the above comparisons of test data with corrosion are based on what are probably the most detailed and specific data available, they are not entirely satisfactory because of the lack of precision of all the classes of data involved. Other comparisons of soil resistivity with pit depths on pipe lines have been published by Weidner and Davis [112], Fitzgerald [113], Gill and Rogers [114], and Ewing [115]. Putnam [52] has shown the correlation between the results of his soil testing method and the loss of weight and pit depths of the National Bureau of Standards specimens, and Denison [95] has made a similar report on his method.

## XI. DETERMINATION OF THE CONDITION OF A PIPE LINE

# 1. LOCATION OF CORRODING AREAS ON PIPE LINES

Soil tests such as those described in the preceding sections are used to locate places along a right-of-way where corrosion may be expected because of local soil characteristics.

It has been pointed out that the effects of the soil are modified by the way in which a trench is backfilled, by the interaction of adjacent soils, by currents picked up by the pipe line, and possibly by other factors. After a pipe line has been laid, it is advisable to determine not only if it lies in potentially corrosive soil, but also the points where the line is actually corroding. Several methods have been devised for doing this, based on the fact that corrosion is associated with differences of potential and a flow of current.

### (a) MEASUREMENTS OF POTENTIALS AND CURRENT

If the resistivity of a pipe is known, the current in the pipe can be calculated from the difference of potential between two points on the pipe a known distance apart. The measurement of the current in the line from point to point provides data that indicate whether the line is collecting or discharging current. The discharging areas are those at which corrosion is occurring. Contact is usually made by driving steel rods into the earth over the pipe. Care must be taken to avoid or correct for contact resistance. Measurements of pipe-line currents have been described by several authors [97, 116, 117, 118, 119].

Usually the potential values are not reduced to currents. If the measurements are made between equidistant points, the change in value will indicate whether or not the pipe is collecting current. The point of maximum corrosion is usually that toward which current flows from both directions. The currents are seldom more than a few amperes and more often only a fraction of an ampere. Sensitive instruments are, therefore, required to measure the drop of potential along a pipe, especially if the pipe is large, and care must be taken to avoid spurious potentials caused by thermal and other effects. These difficulties can be avoided, at the risk of incurring others, by measuring potential gradients in the earth either over the pipe or in a direction at right angles to it. In such measurements the potentials of the contact electrodes must be con-

sidered as well as the contact resistance. If, as is usually the case, the resistivity of the soil varies from point to point, the effects of this variation must be considered also.

Mudd [120, 121] has reported success with a combination of current and potential measurements. Schlumberger and Leonardon [122] have described a method of locating corrosion by observations at the surface of the earth and have developed and patented apparatus for this purpose. Current flowing to or from pipes can be measured directly by apparatus devised by Pearson [123], Haber [124], and McCollum [125]. These devices are of value chiefly in measuring directly fairly large currents in the earth. All the methods and apparatus, except the last two, measure the current discharged from a considerable area of pipe and disclose only the average current density of the discharge. They, therefore, give data for computing the average rate of loss of weight or the average rate of penetration but do not show directly what the maximum rates are, and it is the maximum rate which determines the time required for the development of a leak.

For small areas exposed to approximately uniform conditions, the ratio of the maximum to the average penetration is in the order of 10 to 1, but for larger areas, such as that of a 20-foot length of 8-inch pipe, the ratio may be much larger. In a line with a good protective coating, all the current may be discharged from a very small area where the coating has been injured. If the anodic and cathodic areas are separated by distances much greater than the distance across which the potential difference is observed, much of the corrosion current will affect the observation. If the anodes and cathodes are close to each other, as on the bottom and sides of the pipe, the total current will not be observed.

In general, the computed value of current will be much less than the total current and, therefore, it will be insufficient to account for the observed corrosion. In addition, the so-called long-line currents have been found to change from time to time, probably because of changes in soil moisture or aeration. Obviously, therefore, no close correlation between corrosion and long-line currents can be observed, and if most of the corrosion results from very local circuits, there will be none at all, as claimed by Gill and Rogers [114], who also concluded that soil resistivity was of no practical value in determining corrosion. These conclusions are not in agreement with those of other equally experienced corrosion engineers [112, 115]. The disagreement may be accounted for partly by the differences in the soils involved, partly by differences in methods and apparatus and partly by the standards by which the data are judged, and by what the engineers may have in mind as alternatives for accomplishing the same purpose.

#### (b) COUPONS

Coupons are small iron or steel plates buried near a pipe line to determine the corrosivity of the soil or, if the coupons are connected to the pipe, the rate of corrosion of the pipe. They are used in a number of ways by different corrosion engineers, and there is a considerable difference of opinion as to their usefulness. Schneider [126] cites a number of cases of close agreement between pipe life and predictions based on the use of coupons.

A number of theoretical objections to the use of coupons have been raised. There may be a difference of potential between the coupon and the pipe because of a gradient in the earth or because of rust or mill scale

on the pipe surface. The soil surrounding the coupon may not be representative of that around the pipe. If current flows to or from the pipe or coupon, the current density will not be the same on the two because there will be a tendency for current to collect on or discharge from the edges and corners of the coupons. If these factors are negligible, attached coupons serve a useful purpose as indicators of the rate of corrosion of a bare or imperfectly protected line. They can also be used to determine whether a line to which they are attached discharges or collects current. However, the pipe as a whole might be cathodic and still have anodic spots, such as the bottoms of pits.

# 2. INSPECTION AS A MEANS OF DETERMINING THE CONDITION OF A PIPE LINE

As it is usually impracticable to examine an entire pipe line periodically, the problem of the number and extent of the examinations necessary to furnish representative data arises. This question was investigated jointly by the American Petroleum Institute and the Interstate Commerce Commission. The Commission concluded that for the purpose of determining the condition of the line for rate-making, inspections of the line at selected places and intervals were unnecessary. The decision was reached partly because the cost of an adequate number of inspections would be excessive and partly because for ratemaking purposes obsolescence was considered to be the controlling factor. In the case of one pipe line [127], an agreement was reached that the life of new wrought iron and steel pipe would be considered to be 331/3 years and that the life would be increased 3 to 7 years by bituminous protective coatings, depending on the number of reinforcing layers. The data upon which the agreement was reached were not published. Obviously, such a basis would not be satisfactory for determining the physical condition of a section of a line in order to decide whether or not it requires reconditioning, protection, or repairs.

Gill [128] concluded, from statistical considerations, that a line should be inspected at equally spaced points, and that 2,000 feet was the greatest permissible distance between inspection points. Such inspections convey an idea of the average condition of the line, but only detailed inspection of the data will reveal where corrosion is in progress.

Logan and Koenig [129] made statistical studies of pit depths on pipe lines, including one 200-mile length on which the deepest pit on each joint of pipe was measured. They attempted to show the different factors affecting the results of inspections, including the space interval between the sections inspected, the number of inspections, the location of the starting point from which equally spaced inspections were made, and the effect of the size of the area inspected on the observed average maximum pit depth.

Each of the first eight columns in figure 67 represents the average depth of the deepest pit on each 25 joints (20 feet long) of 8-inch pipe. The inspections were made at 1-mile intervals. The different columns represent the data for eight different starting points separated by ½-mile intervals. In two cases the computed average differs from the true average by about twice the standard error, whereas, for normal data, the probability of such an occurrence is but 1 in 20. It is evident that for a comparatively few inspections the starting point may affect the data. The next four columns in figure 67 show the results of 4 sets of 50 inspections of the same line. None of the computed averages differ

from the true average by more than the standard error which, of course, is smaller than that for 25 observations. The last two columns in the figure indicate that the reliability of the average is only slightly increased by increasing the number of inspections to 110, i.e., making the inspections at ¼-mile intervals. This conclusion applies strictly, of course, only to the section of the line under consideration.

Figure 68 shows the effect of the area inspected and of the starting point for 30 inspections of a line at 1-mile intervals. As the area inspected is increased, the effect of the starting point is decreased, as is also the standard error of the average, even though the number of

inspections is constant.

The relation between the inspection interval, starting point, and number of inspections when the corrosion criterion is the condition of the pipe is illustrated in figure 69, supplied by the Stanolind Pipe Line Co. The figure indicates, as did figure 67, that if the number of inspections is small, the average is affected by the starting point of the inspections.

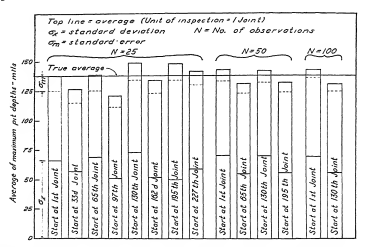
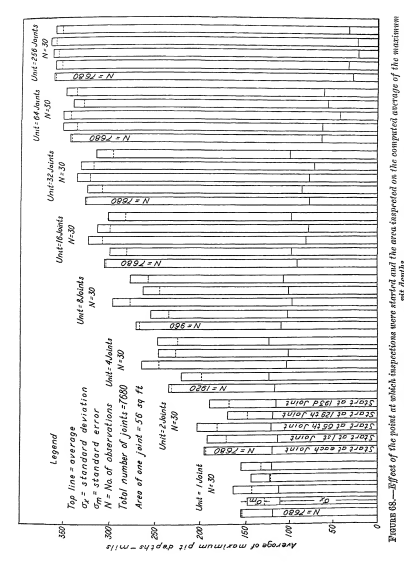


FIGURE 67.—Effect of the starting point and the number of inspections on the observed average of maximum pit depths on sections of a pipe line.



pit depths.
Note that the value of the pit depths increases as the area inspected increases.

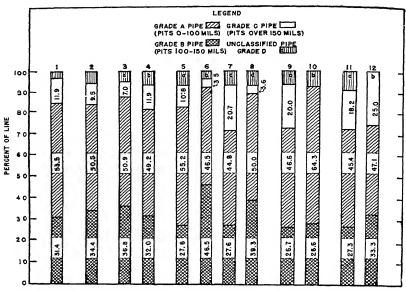


FIGURE 69.—Effect of method of inspection on apparent condition of a pipe line.

(2) ... 500 feet (3 and 4) ... 1.000 feet (5 to 8) ... 2.000 feet (9 and 10) ... 4.000 feet (11 and 12) ... 5,000 feet Inspection data:
11.4 miles of pipe line,
Laid in 1914, taken up in 1934.
One 20-foot joint of pipe is inspected at
each inspection point.

The relation between the size of the area inspected and the depth of the deepest pit and the effect of the area on the reproducibility of the data are important considerations in determining how much pipe should be exposed at each inspection point and the frequency of inspection (fig. 70).

The lower curve is based on data from a 9-year-old 12-inch steel line exposed to a heavy clay soil, assuming that an inspector started at 48 different points on the line and first uncovered 1 foot of pipe and measured the depth of the deepest pit. The trench was then extended 1 foot and the depth of the deepest pit on the 2-foot section was measured. Similarly, the depth of the deepest pits on greater lengths up to 20 feet were measured. The curve indicates that the average depths of the deepest pits increase as the area inspected is increased, but that when a large number of observations (48) is made, the standard error of the average (one-fourth of the length of the vertical lines) does not decrease very much. The average of the values obtained by a large number of inspections of 1-foot lengths is almost as representative as the averages of the values obtained on entire joints of pipe. This conclusion would not hold if only a few inspections of 1-foot lengths of pipe were made, as the pit depth on a length of pipe varies greatly from foot to foot. The upper curve indicates that each inspection point would have to include a large number of lengths of pipe before the value of the average maximum pit depth would become substantially constant. The curve also indicates

that when this area is reached, the number of observations required for a specified degree of reproductibility decreases as the area examined is increased.

Logan and Koenig suggested that, since corrosion appears to be characteristic of the type of soil to which the pipe is exposed, and since the same type may occur in several places along a long right-of-way, the number of inspections necessary for determining the condition of the line might be reduced by first identifying the soils traversed by the line and then making in each soil only a sufficient number of inspections to establish its corrosiveness. In addition to reducing the number of inspections, this procedure would furnish data as to the location of the sections which need reconditioning or protection. The relation of soil types to corrosion has already been discussed and is illustrated in figures 50 and 51.

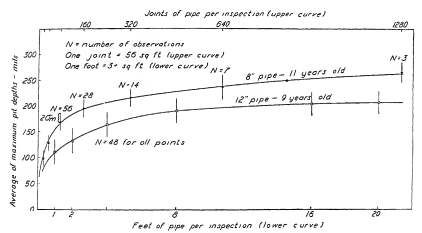


Figure 70.—Relation of area of pipe inspected to the maximum pit depths on each area.

All investigators seem to agree that a large number of inspections is esssential to an accurate determination of the condition of the pipe. If the number is adequate, the size of the area is unimportant, but the number of inspections required for a certain precision of the results will increase as the area inspected is decreased.

The conclusion that pipe should be inspected at equally spaced distances or with regard to soil types is based on the assumption that the condition of the entire line is desired, as for valuation purposes. If the operator of the line has had considerable experience with corrosion and is interested only in anticipating leaks, he may follow the practice of the Susquehanna Pipe Line Co. described by Van deWater [130]. This organization inspects its pipe at places where abnormal corrosion is likely, such as at low spots, bogs, swamps, and highway and railroad crossings. Locations where a leak would be particularly dangerous are also examined. Thousands of places on a line extending through Pennsylvania and Ohio have been thus inspected. From 7 to 10 feet of the pipe are exposed at each inspection point, and

from 10 to 15 pit depths are measured. Examinations cost about \$10.00 each and average about \$65.00 per mile of line. About 10 percent of the line has been protected. This is accomplished by boxing the pipe and filling the box with asphalt. The protected sections range from 20 to several hundred feet in length.

## XII. TESTS OF COATINGS

Various tests have been devised for determining the suitability of a coating for a given kind of service or its condition either before or after exposure to corrosive soils. In general, they are used to determine thickness, continuity, electric resistance, or mechanical strength.

### 1. METALLIC COATINGS

The use of metallic coatings underground is not general enough to warrant an extended discussion of methods of test of such coatings. The effectiveness of a metallic coating depends in part on its thickness, and this factor should be taken into consideration in comparing or specifying metallic coatings for pipes or tanks. The potential of the coating with respect to the underlying metal is also important because galvanic corrosion may occur if both metals are exposed to the soil.

Burns and Schuh [30] have described a number of tests that may be classified as (1) stripping tests, in which the loss of weight of the coated metal is determined after removing the coating, (2) solution tests, in which the time required for rust to appear is determined when the coating is subjected to the action of certain corrosive solutions or sprays, (3) electrolytic tests, in which current is passed from the coated metal to a paper in which the anion from the underlying metal is precipitated. (4) the chord test, in which the coating is cut at a definite angle and its thickness measured with a microscope. These tests, with the exception of the electrolytic test, are destructive, and their usefulness therefore is limited mainly to investigational work.

### 2. INSULATING COATINGS

The principal causes of failures in insulating coatings are faulty application, injuries, pinholes, decay, and distortion resulting from what Scott [131] calls soil stress.

Tests for continuity and resistance have been developed by Ewing and Scott [132]. The section to be tested is first washed and then painted with a slurry of kaolin or iron-free clay and water to fill holes, cracks, and depressions. A sheet of absorbent paper known to plate printers as wiping paper and sometimes sold as pinhole paper is placed over the clay and is wrapped with several layers of water-soaked cotton flannel 1 foot wide. The pad is held in place by a saddle consisting of a framework of copper strips, as shown in figure 71. A 22-volt battery is connected between the saddle and the pipe, the pipe being made the anode. The circuit is closed until the product of the applied voltage and the time in minutes is approximately 100. The paper is then removed and washed in a 1-percent solution of potassium ferricyanide. current flows from the pipe through openings in the coating, ferrous exide will be deposited in the paper, which will cause blue spots to appear corresponding to the holes in the coating upon treatment with potassium ferricyanide. As the number and size of the spots are indications of the condition of the coating, this test is known as the pattern test for coating

continuity. The pattern test is helpful in locating small pinholes and cracks; patterns from different coatings, however, are not comparable unless the resistivities of the waters used and the voltage-time products are the same for each coating.

If the current and voltage supplied by the battery as used in the pattern test are observed, the resistance or conductance of the coating can be calculated. The latter is usually expressed as micromhos per square foot. Although the conductance measured in this way involves the conductances of the pad, paper, and test solution, these are usually negligible if the coating is good; there is nothing to be gained by testing a coating with an obvious hole. As the range of conductance values of coatings is very large, a special instrument, such as that designed by Shepard [74] for measuring them, is desirable.

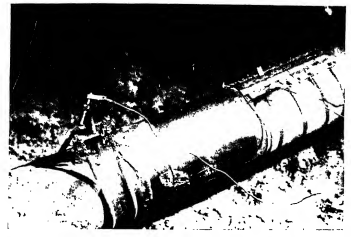


Figure 71.—Setup for making pattern and resistance tests of pipe coatings.

The pattern and conductance tests are, of course, applicable only to small areas of coatings after they have been exposed to soil or water long enough to permit the moisture to pass through any holes in the coating. There are two other types of continuity tests that are applicable to coated lines and which can be used to determine the condition of large areas of coatings. One of these is a high-voltage or spark test intended primarily for the detection of pinholes.

The apparatus, figure 72, consists of a high-voltage transformer, or induction coil such as a Ford spark coil, a battery, a current interrupter, and an insulated metal brush. One side of the secondary of the coil is attached to the pipe and the other side to the brush. The battery and interrupter are connected across the primary of the coil. As the brush is passed over the surface of the coating a spark will jump to the pipe when the brush is over a pinhole or holiday; the sound and flash of the sparks thus locating the flaws in the coating. Clarvoe [133] and Harrell [134] have described such apparatus and its use. These detectors have several faults. The operator may skip some of the coating area. The high voltage may break down the coating if it

is applied too long at one spot. The voltage may be too low if the coating is very thick. Most forms of the apparatus do not locate thin spots in the coating. If the output of the transformer is high, accidental contact with the secondary circuit may cause a serious accident. Such holiday detectors have been used extensively and have been instrumental in the production of better coatings. Their use is specified in many contracts for coatings. Recently, Stearns [135, 136] patented an improved form of detector that travels over the pipe in such a way that the missing of a pinhole is unlikely. The apparatus operates a counter, lights a colored light, and rings a bell whenever a spark jumps to the pipe. The voltage of the apparatus can be controlled. After a spark has passed, the high voltage is cut off for 1 second.

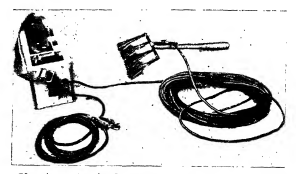


FIGURE 72.—Apparatus for locating pinholes in bituminous coatings.

It consists of a Ford spark coil, dry batteries, and a metal brush with an insulated handle.

Although the use of a holiday detector may insure the application of a continuous coating, it does not insure a continuously coated pipe, because the coating may be injured after the test especially in the course of laying the coated pipe. In addition, the coating may be punctured by roots, as illustrated in figure 73, or it may crack after it is in the ground. The apparatus described by Pearson [137] is therefore of interest. A 1,000-cycle battery-driven hummer is connected between the coated pipe and a rod driven into the ground a 100 feet away. Most of the current that passes from the pipe to the ground rod must flow through holes in the coating. This results in an increased IR drop of potential in the soil opposite the holes. The disturbed earth potential is located by two men 20 feet apart, one of whom carries an amplifier and indicating meter, the other man providing the ground connection. Each man has steel cleats on his shoes, connected in the first case to the amplifier shielding and in the second case to the amplifier input by means of a shielded cable. They walk along the pipe line as nearly over its center as possible. When either man passes over a hole or poorly insulating spot in the coating there is an increase in the deflection of the indicating meter and in the sound in the earphones in parallel with the meter. Although the use of the apparatus requires some experience, its effectiveness has been demonstrated by several users. Pearson [137] has published details concerning the design and use of the apparatus. It will locate individual pinholes and other openings in an otherwise good coating or areas of poor coatings if pinholes or carbonized particles are

too numerous to make a single defect outstanding. In this article Pearson also describes a method for measuring the resistance of a coating

on a pipe line without uncovering the line.

In order to compare the ability of different coatings to withstand soil stress, Ewing [109] constructed an outdoor soil box in which the conditions producing the stress could be simulated. The box, about 1





FIGURE 73.—Coatings injured by roots.

4. Root beneath a rag-felt-reinforced asphalt coating; B, grass roots in and beneath a 10-year-old asbestos-felt shielded coal-tar enamel coating on a pipe line in Oklahoma.

foot in depth was constructed of porous brick supported above the ground to provide the greatest possible surface for evaporation. The specimens under investigation were first subjected to the pattern test and then buried in the box so that there was about 6 inches of dry soil above the specimens. Since the available soil did not have a sufficient

volume shrinkage between saturation and complete dryness, it was modified by the addition of about 1 percent by weight of sodium carbonate (washing soda) which increased the shrinkage from 28 percent to about 53 percent. By alternately wetting the soil and letting it dry, the samples were subjected to a number of cycles of stress and then examined. Soft coatings were found to be distorted, but not so much as some specimens observed in field tests, probably because the soil in shrinking tended to pull away from the sides of the box instead of from the coating. There is a difference of opinion as to whether the distortion results from a pull when the soil shrinks or from pressure when the soil swells as it absorbs moisture.

Ewing also tried impact tests by dropping a steel ball weighing 1.65 pounds from various heights and noting the minimum height at which a failure was produced as indicated by an electrical test. He found that the test, as he applied it, involved several variable factors difficut to control.

# 3. TESTS OF COATING REINFORCEMENTS AND SHIELDS

To reduce distortion of bituminous coatings by clod pressure, pipe movement, and soil stress, the bitumen is frequently reinforced by a spiral wrapping of fabric, which may be an open-mesh woven cotton

-				
No.	Description of material	Weight	Thickness	Ratio of strength of bottom half of specimens after exposure of 301 days to original strength ¹
1 2 3 4 5	14-lb. asbestos felt, asphalt-saturated. 15-lb. asbestos felt, asphalt-saturated. 15-lb. asbestos felt, tar-saturated. 14-lb. asbestos felt, tar-saturated. Same as 1 with bakelite resin saturant.	17.0 14.6	Mils 26.7 28.1 29.5 27.4 25.9	0.88 .87 .49 .745
8 9 10	15-lb. asbestos felt, tar-saturated 15-lb. asbestos felt, grease-saturated 15-lb. rag felt, asphalt-saturated 30-lb. rag felt, asphalt-saturated 30-lb. coated rag felt, asphalt-saturated	15.8	28.7 28.6 41.6 69.8 55.2	.93 .445 .21 .22 .25
11 12 13 14 15	40-lb. coated rag felt, asphalt-saturated 15-lb. rag felt, tar-saturated 30-lb. rag felt, tar-saturated Polymerized resin shield, Polymerized resin shield, paper-backed	15.5	96.8 40.4 65.2 10.5 19.7	.26 .00 .07 .12 .105
16 17 18 19 20	Cotton fabric with rot inhibitor, grease-saturated. Cotton fabric without rot inhibitor, grease-saturated. Woven asbestos fabric, open weave, tar-saturated. Cotton fabric, asphalt-saturated. Cotton fabric, tar-saturated.	8.7 8.0 21.5	21.0 18.5 73.0 32.7 33.7	.11 .0 1.37 0.00
21 22 23 24 25	Coir fiber, close weave. Burlapp Manila fiber Manila fiber. Manila fiber, creosote-treated. Hemp	22.0 7.8	97.7	.00 .00 .00 .16
26	Sisal	48.6	214.7	.00

 $^{^{1}\}mathrm{Based}$  on average strength after soaking. The measurements of February 1934 and January 1935 were used.

fabric of the Osnaburg type, burlap or bitumen-saturated rag, or asbestos felt. This wrapping may be covered with bitumen, in which case it is called a reinforcement, or it may be left exposed directly to the soil, in which case it is called a shield.

To determine the effect of soil, or rather of soil bacteria, on these materials, Scott and Ewing [138] exposed 26 varieties of these materials to a soil obtained from the flood plain of a river near Washington. Strips of the materials were buried in boxes of the soil, which was saturated at intervals of 1 month. Table 99 shows the character of the materials tested and the strength of the materials after 301 days of exposure in terms of their original strengths. It indicates that all the organic materials deteriorate when exposed to soil bacteria. The materials were exposed directly to the soil, and some were not impregnated with bitumen or treated to resist rot. In service, they might be saturated with bitumen and covered by a fairly thick layer of this material. Under these conditions, they should last much longer.

When Ewing [139] removed the first set of AGA specimens of coatings, he desaturated the fabrics and then determined their strength. He observed little or no deterioration of the fabrics saturated with coal tar after 9 months of exposure to organic soils, but the organic fabrics in asphalt coatings had deteriorated. He found also that the rotting decreased with increased thickness of the asphalt over the fabric, and that

muck soils were the most destructive to the fabrics.

### XIII. PREVENTION OF CORROSION

The investigations reported in Section XII were carried out to obtain data on the relative corrosiveness of various soils, the relative resistance to corrosion of various materials commonly used underground, the influence of various factors other than the inherent corrosiveness of a given soil on the rate of corrosion of a material buried in it and to develop methods preventing or minimizing losses due to corrosion. The protection to use in a given case is dictated by economic considerations and depends upon a variety of factors. If obsolescence is an important factor, the added expense involved in protecting a pipe line so that it would last a very long time might be unjustified. In any event, the increased cost of any protective measure which might be adopted, including the annual charges on the increased cost, should be balanced against the cost of replacement or repairs. Special protection may be unnecessary or unduly expensive in soils that are only mildly corrosive. On the other hand, more severe conditions may require some protective measures, such as the use of corrosion-resisting materials, special treatment of the soil, protective coatings or cathodic protection.

# 1. MATERIALS SUITABLE FOR USE IN MILDLY CORROSIVE SOILS

According to the data presented earlier, the different ferrous alloys commonly used in pipe corrode at approximately the same rate. In some soils wrought iron has a tendency to corrode somewhat more uniformly or to show a slightly lower average penetration than other metals. In a few alkaline soils, cast iron appears to have a somewhat higher rate of penetration, but this may be compensated for to some extent by the strength of the corrosion products. The differences, however, are not very great. From this, it would appear that, at least in mildly corrosive soils in which bare pipe would ordinarily be laid, the thickness of the pipe wall is more important than the material of which the pipe is made.

If a pipe line is to be laid in a soil that is only mildly corrosive, the length of service will depend primarily upon the wall thickness of the

pipe. In some soils (table 51) a light-weight steel pipe will render many years of service without a leak. Under such conditions, a pipe only heavy enough to withstand the internal and external pressures with a reasonable factor of safety may prove to be the most economical.

The pit-depth-time factors (table 51) indicate that a moderate increase in wall thickness of a pipe may be expected to greatly increase its useful life in many soils. Figure 16 and curves plotted from the pit-depth-time equation show the required wall thickness for a given average pipe life before puncture and indicate that in many soils it is practicable to obtain long service life by means of a moderately thick pipe wall. Such curves are based on average values and considerably thicker pipe walls should be used if entire freedom from punctures is desired. It may not be economical, however, to use a very thick pipe in order to secure a long life. Obsolescence and the annual charges on the increased cost of the longer-lived pipe may outweigh the advantage of the longer service.

## 2. CORROSION-RESISTANT MATERIALS

Various attempts have been made to reduce losses due to corrosion in the more corrosive soils by the use of noncorrodible or corrosion-resistant materials. Many early water pipes were made of board logs and, later, of pipes made of wood staves held in place by metal bands. If kept con-

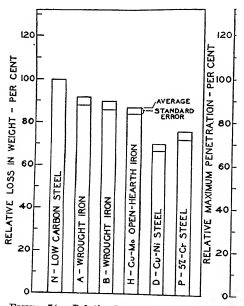


FIGURE 74.—Relative loss of weight and penetration of wrought ferrous specimens.

Low carbon steel=100 percent.

tinuously wet, these pipes often last for many years, but under some conditions the wood-stave pipes leak badly and the metal bands corrode and break.

Cement and concrete, usually reinforced with steel, have been em-

ployed extensively for water, especially for large volumes at low pressures. For many years in Europe, and more recently in the United States, pipes made from a mixture of cement and asbestos have been used. Some data on this type of pipe were given earlier (page 128), but they do not cover a sufficient period of exposure to warrant definite conclusions, although they appear to be satisfactory from the corrosion standpoint. For sewers and drains, virtified tile is satisfactory as far as corrosion is concerned.

In most soils, copper and copper-rich alloys corrode much more slowly than iron or steel and the corrosion is more uniform. There are not enough data to indicate definitely how the rate of corrosion in copper and copper alloys varies with time, but apparently the rate of corrosion becomes nearly constant after a short time if the soil conditions do not change. If copper or copper alloys are used in conjunction with ferrous materials, they should be connected by insulating joints to prevent accelerated corrosion by galvanic action. Muntz metal (60% Cu, 40% Zn) is unsuitable for service in many soils on account of dezincification, but

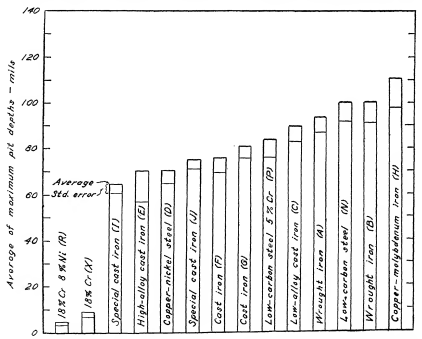


Figure 75.—Average of maximum pit depths on ferrous pipes removed from Merced clay adobe after an exposure of 5 years.

corrodes less rapidly in tidal marshes, even though it is subject to some dezincification. Lead pits badly in a few soils, but corrodes little in others, especially those containing sulfates.

Ferrous metals can be made more resistant to corrosion by the addition of certain alloying elements, such as copper, chromium, or nickel.

It has been found, however, that small additions of these elements are not effective for underground service. This is shown in figure 74. Steels containing appreciable percentages of chromium and nickel corrode very little underground (fig. 75). Except in a relatively few soils, cast iron containing 15 percent of silicon does not corrode appreciably but unfortunately this alloy cannot be easily machined or welded. The use of ply metal, that is, corrodible metal surrounded by a thin layer of corrosion-resistant material, such as copper or stainless steel, has been suggested. One such material was included in the National Bureau of Standards tests, but little or none of this material is in service underground. Copper-clad steel is quite generally used for ground rods and similar purposes. As suitable corrosion-resistant materials are expensive, it is usually more economical to employ other means for reducing losses due to corrosion, with the exception perhaps in the case of small-diameter pipe or when corrosion might result either directly or indirectly in serious losses.

### 3. SOIL TREATMENT

Soil treatment may consist in (1) the addition of chemicals to neutralize the corrosive properties of the soil or to accelerate the formation of protective films, (2) the replacement of corrosive soil next to the pipe

by less corrosive soil, or (3) drainage.

Attempts to prevent corrosion by the neutralization of the corrosive properties of the soil are based largely on a misunderstanding of the chief cause of underground corrosion. Only a very few soils are chemically corrosive. To be effective, an added chemical must be in solution and in this condition it will be rapidly removed from the neighborhood of the pipe by diffusion and the movement of water in the soil. Hydrated lime and ground limestone are the chemicals most commonly used, because they are moderate in price and only slightly soluble. To minimize losses by diffusion, Grodsky [140] proposed to surround the pipe with a waterproof membrane and place soil mixed with sufficient lime or other agent to neutralize its acidity between the membrane and the pipe.

The use of lime has been stimulated by the observation that, frequently, pipes in soils containing lime are coated with a thin scale of calcium carbonate and that the pipe beneath this scale is not seriously corroded. There appears to be little available information as to the extent of the use of lime to neutralize soil in trenches or of the success of such use. The Dutch Central Corrosion Committee [6] reports the examination of a 9-year-old pipe laid in a bog. The pipe was surrounded by a layer of sand to which lime had been added. Analysis of the sand showed that it contained 5.1 percent of CaCO₃ and had a pH of 7.9. The pipe was not corroded, except at a point where the sand was no

longer in contact with the trench.

Wichers [141] made extensive studies of underground corrosion in Groningen, Holland, and suggested three methods of improving soil conditions: (1) using impermeable earth for the upper layers of the trench and drainage where necessary, (2) kneading plastic clay or loam around the pipe, and (3) surrounding the pipe with sand or earth neutralized with lime. Clay and sand have been used to surround short stretches of pipes in the United States, especially when the pipe passed through made ground containing cinders. The practice is not general and no definite results have been reported but no doubt these materials are less corrosive than cinders and rubbish. More often a heavy bituminous coating or one of concrete is used for such conditions.

# 4. PROTECTIVE COATINGS

The most common method of reducing losses due to corrosion underground is by the application of some form of protective coating. The coatings ordinarily used on the larger pipes are nonmetallic, but zinccoated (galvanized) pipe is extensively used in the smaller sizes, especially for water services.

Kelly [142] states that approximately half of the steel pipe currently in service is protected by some form of coating, and that 80 percent of such pipe is covered with a tar-base coating. Excepting the dip coating applied to nearly all cast-iron water mains, the most commonly used coal-tar coating is an enamel made by adding finely divided inert material to coal-tar pitch. Coatings of concrete or grease are also used to a certain extent.

### (a) CONCRETE

Concrete and cement-motar coatings have been used extensively by certain pipe line companies when corrosion is very severe. The earlier cement coatings were applied after the pipes were in the trenches by placing boards 1 to 2 inches from the pipe and filling with mortar. Sometimes no board was placed beneath the pipe, and occasionally mud was forced ahead of the mortar and into contact with the pipe. This ultimately resulted in leaks, and in some cases, the concrete cracked longitudinally. One company has [143] developed a metal form which is put around the pipe above the trench. This is filled with concrete and removed a day later, when the concrete has set. This method makes it possible to use a relatively thin coating, thus reducing the cost. Concrete applied to pipe by means of steel forms was tested at site V of the API coating tests. Twenty feet of pipe beneath this coating was examined after 10 years. The maximum pit depth was 24 mils and the average on each 1 foot of pipe, 5 mils. The maximum pit depth on 10 feet of adjacent bare pipe was 50 mils and the average 38 mils. The 10-year-old API coatings examined in 1940 included 170 feet of asphalt emulsion shielded by cement mortar having an average thickness of 0.419 inch. Sixty-one 1-foot sections of the pipe beneath this coating were pitted. The depth of the deepest pit was 107 mils. It has been stated that the alkali in the concrete makes the pipe beneath it cathodic with respect to adjacent uncoated pipe and, therefore, accelerates the corrosion of the bare pipe.

The deepest pit on any of the uncoated pipe at site V was 65 mils and the average of the deepest pits on 13 uncoated 8-foot sections was 49 mils. These data do not support the theory that the coating accelerates the corrosion of adjacent bare pipe. If such a phenomenon should exist, it could be overcome by the use of insulating joints at the ends of the coated sections or the application of an insulating coating, such as coaltar enamel, to a few feet of the line adjacent to the cement coating. Another solution would be the application of the coating to the entire line. It will be noted that the concrete was not sufficiently alkaline to prevent corrosion of the metal beneath it.

Occasionally, a condition occurs where it is not convenient to remove the pipe from the trench for the application of a bituminous coating or where the weather is such that it is difficult to keep the pipe dry over a sufficient period to permit the application of a hot bituminous coating. In such a case, or if local labor unfamiliar with handling bituminous coatings has to be employed, concrete may be the best coating to use.

### (b) GREASE COATINGS

Grease with or without an added inhibitor, such as a chromate, has been used extensively by several pipe line operators. Usually the grease is protected by a coat of wax or by a wrapper of fabric or metal foil. Greases have the advantages that they can be applied cold by hand to a pipe which is not dry and that they tend to flow and so to heal any small openings in the coating. They have been used successfully under wet soil conditions. Bredberg [144] has written an interesting description of such an application of grease.

### (c) BITUMINOUS COATINGS

The various kinds of bituminous coatings can be divided into two general groups—asphalt base and coal tar base. Each base has certain characteristics that make it preferable to the other, but each has some characteristics that are objectionable. Within each group, the materials vary so greatly that it cannot be said that one base material is superior to the other. In general, the asphalt-base materials are less susceptible to shock and changes in temperature, but they tend to change more and to absorb or transmit more moisture than the coaltar-base materials. The so-called plasticised coal-tar enamels are more nearly like the asphalts in their response to changes in temperature, but the range of temperatures within which they can be successfully applied is smaller than that of the older enamels.

The fundamental requirements of a satisfactory bituminous coating are permanence, continuity, and low electric conductance, but certain other characteristics are necessary for a satisfactory coating. The hardness or softening point, as measured by standard ASTM methods, should be as high as is consistent with freedom from cracking and satisfactory application, and this will depend on the character of the bitumen, the method of application, and the season and the locality where the coating is applied or used. Continuity depends mostly on the care with which the coating is applied to the pipe, but ultimately it will be affected by the resistance of the coating to cracking and cold flow under pressure.

Many bituminous coatings have a high coefficient of expansion and should be protected against large rapid changes in temperature. High dielectric strength, that is, ability to withstand high electric potentials, is not in itself important, but the coating should have a low conductance or high resistance to low potentials. The fact that a coating when applied to a pipe will withstand several thousand volts has little significance except as an indication that the area tested is free from pinholes. Measurement of electric resistance after exposure to water for several months is a better test.

Adhesion of the coating to the pipe has often been stressed. Its importance lies in the fact that if the coating does not adhere, water entering at one point will spread beneath the coating, and the resulting corrosion may lift the coating and expose still more pipe to corrosion.

Probably the best way to secure continuity and long life for a bituminous coating is to use multiple applications. If a reinforcement is used to obtain thickness, it should be of inorganic material. It has been shown that organic fabrics exposed to moisture may rot. A stiff permanent wrapper will distribute the pressure on the coating and reduce distortion. A light bitumen-saturated asbestos-felt wrapper is not sufficiently rigid to prevent distortion of soft bituminous coating and may be penetrated by grass roots. Figure 73A, shows a root that has penetrated an asphalt coating reinforced by two layers of rag felt. Figure 73B, shows grass

roots in an asbestos-felt wrapper applied to a coal-tar enamel. These figures are of interest because they indicate that sufficient moisture to permit plant growth penetrated the bitumen-saturated wrappers. Figure 3 shows the distortion of a rag-felt-reinforced asphalt coating exposed to a heavy clay soil that cracked on drying. The figure also shows cracks in the coating, which may be the result of soil movement or of deterioration of the bitumen. Figure 76 shows a cotton-fabric-reinforced-asphalt coating that cracked badly, probably because of evaporation of volatile constitutents. In the same figure is shown a coal-tar-pitch coating that flowed from the top of the pipe. The resulting corrosion was much more severe than on nearby uncoated pipe.



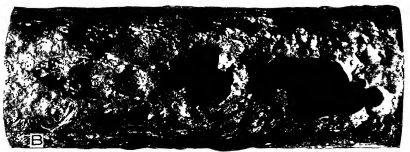


FIGURE 76.—Examples of failure of pipe coatings.

A, Cotton fabric reinforced asphalt coating—note cracks; B, a coal-tar coated pipe from which the coatings flowed because it was too soft. Soil—tidal marsh.

Any specification for bituminous coatings should include a high-voltage test of the bitumen before reinforcement or wrapper is applied to the coating, if this is practicable. Care should be taken that every part of the coating is brought into intimate contact with the brush or other terminal of the high-voltage circuit. If the pipe has been transported after coating, it is advisable to test the coating again just before it is placed in the trench. Great care should be taken to see that the bottom of the trench is free from stones and foreign material and that such materials are not allowed to come in contact with the coating when the trench is back-filled. Care should also be taken to see that the coating is not injured by skids, chains, or other tools used in laying the pipe, as faulty application and accidental injuries are the most common causes of coating failures.

An inspection of the coating by means of the Pearson [137] apparatus a year or so after the pipe is laid will show whether or not the coating

has been properly applied and handled. Subsequent inspections by the same method will indicate the progress of coating deterioration, if any.

The American Water Works Association [145] has prepared specifications for coal-tar enamels for use under different conditions. They are not suitable, however, for coal-tar enamels of the older types or for asphalt-base coatings. The tests include softening point, percentage and fineness of filler, specific gravity, penetration, high-temperature, low-temperature, and impact tests. Only experience will show whether or not the use of these specifications will insure a good coating material, but they are certainly a step in the right direction.

It frequently has been said that as a result of the API tests great improvements in pipe coatings were made. It is of interest, therefore, to consider the type of coating selected by the War Emergency Pipe Line [146], an organization composed of prominent pipe line operators, for the most important pipe line in the United States. Most of this line received a coat of coal-tar primer followed by a coat of coal-tar enamel and a wrapper or shield of asbestos felt. The coatings were machine applied at the right-of-way and have an average thickness of  $\frac{3}{3}$ , inch. Most of the line was not electrically tested for pinholes. The reason for the choice of this type of coating is not entirely clear, but it may have been influenced by the exigencies of the situation—availability of materials and machines, ease of application, etc. Whether or not the choice was a wise one will not be known for a decade or more. The Products Line [147] constructed by the same organization, was similarly protected. However, it was triple-coated and wrapped at river crossings. In the wet, corrosive soils of the Gulf Coast area, some of the feeder lines were coated with an asphalt mastic coating similar to coating O in the API tests but which was applied by a somewhat different machine, which extruded the mastic around the pipe without seams.

As previously noted, one company protects its 6-inch gasoline lines by enclosing the sections exposed to corrosive soils in a cypress box with a 2-inch space between its sides and the pipe. The pipe is supported by porcelain blocks. The box is lined with roofing paper and filled with asphalt. The expenses of this treatment of short sections of pipe is thought to be less than cathodic protection, and is preferable because less attention is required.

Another large pipe line operator [148] uses a petroleum asphalt applied at the site by machine and shielded by a spiral wrap of asbestos felt. The thickness of the coating is 32 inch. Sterling [148] reported that after 13 years a pipe coated with asphalt and asbestos felt was in excellent condition, whereas unwrapped sections of this pipe were almost beyond recovery, many joints having from 15 to 18 penetrations.

Thomas [149], who for many years has taken a leading part in the development of the pipe coatings in Australia, reports that the following coating has been adopted for the protection of large-diameter water pipes in Melbourne:

Pitch, vertical retort, gasworks; softening point	
170° to 180° F	34%
Tar, horizontal retort, gasworks, crude	26%
Limestone (200 mesh)	20%
Steatite (200 mesh)	14%
Aspestine	6%
Melting point of enamel, ring and ball 175° F	0,0
Working temperature for application, 450° to 500° F.	

It may be noted that the percentage of filler is higher than that ordinarily used in the United States and that much of it is limestone. Most specifications in this country require an inert filler, and some prohibit the use of calcium carbonate and slacked lime. Rag felt is used as a shield because asbestos felt is not manufactured in Australia.

Elaborate apparatus has been developed for the application of the enamel to the inside and outside of pipes. For the outside coating, two layers of enamel are applied after which they are spark tested, a layer of rag felt is then applied and flooded with a third coat of enamel. Finally, the coating is whitewashed. The thickness of the coating is 0.24

inch. The interior coating is spun on to a thickness of 0.1 inch.

Apparently, asphalt is preferred in Holland, as the Dutch Corrosion Committee [3] has issued detailed specifications for the testing and application of asphalt coatings in several thicknesses. Kieselguhr, clay, gypsum, or other water-soluble salts must not be used as fillers. Asphalt-saturated wool felt and impregnated jute are specified as reinforcing materials. Four degrees of protection are specified—

1. Light protection (not to be applied to steel pipes):

A. A coat of blown asphalt.

B. A wrapping of waterproof paper to protect the coating against damage during transport and laying.

2. Moderate protection:

A. Priming coat of blown asphalt.

B. Topcoat of filled asphalt with or without a protective wrapper.

3. Heavy protection:

A. Priming coat.

B. Topcoat of filled asphalt.

C. Reinforcement—jute or wool felt.
D. Protective coat of filled asphalt.

4. Very heavy protection: A. Priming coat.

B. Topcoat of filled asphalt.

C. Reinforcement; impregnated coir cloth or impregnated asbestos felt.

D. Protective coat of filled asphalt.

E. Paper wrapping.

F. Straw cord wrapping.

Little information has appeared as to the effectiveness of the newer pipe coatings, partly because a decade or so must elapse under most soil conditions before corrosion becomes serious even on poorly coated pipe, and partly because sponsors of coatings are averse to the publication of results that do not meet their expectations. Furthermore, the shortage of manpower has prevented any inspections of coatings that could be postponed.

Rogers [150] has reported some data on new enamel coatings. One coating consisting of an enamel and asbestos-felt wrapper showed a resistance of from 400 to 9,000 ohms per linear foot of pipe after 2 years, and changed little during the next 2 years. A second machine-applied enamel shielded with asbestos felt and applied in 1940 had an initial resistance of 67,500 ohms per foot. After 14 months the resistance ranged between 375 and 11,000 ohms per foot. Another machine-applied enamel showed nearly a uniform resistance of 6,200 ohms per foot after 14½ months. The same materials, hand-applied to another section of line, showed resistances ranging between 3,875 and about 200 ohms per foot

after 15 months of exposure. Three interesting facts are shown by the data. During the first few months after burial one 4,000 foot section of one coating dropped from a very high value of resistance to a minimum of 400 ohms per foot. After the initial drop in resistance the coating showed little change in resistance. The resistance of the coatings

differed greatly at different sections of the line.

Similar data were reported by Hadley [151]. A section of an 8-inch steel line laid in a dumping ground was protected by three coats of coaltar enamel, shielded by wrappers of asbestos felt and kraft paper, machine applied. The thickness of the coating was 5/16 inch. The line was divided into 2,000-foot sections by insulating joints. Extreme precautions were taken to insure a perfect coating and high insulating joint resistance. To prevent injury by the back-fill, the pipe was surrounded by a layer of sand. Test wires were brought out from each section of the pipe. When installed the average resistance of the coating on 14,300 feet of the pipe was 94,000 ohms per linear foot. At the end of 3.8 years the average resistance of the same sections was 9,300 ohms per foot. Initially, the maximum resistance of one 2,000 foot section was 429,000 chms per foot. The final resistance of this section was 2,600 ohms per foot. One section had a resistance of 536 ohms per foot after 3.8 years. Most of the changes in resistance occurred between the third and the ninth month of exposure.

At the 1944 convention of the National Association of Corrosion Engineers, Hugo [152], in describing the use of the Pearson coating-tester, reported that an inspection of about 150 miles of an asphalt mastic coating, most of which was about 1 year old but with 11 miles of 3%-year-old coating, disclosed 80 electrical imperfections. Eleven of these were examined and five obvious coating injuries were found. At six points the imperfections were not visible. A similar test of 45 miles of asbestosfelt-shielded coal-tar-enamel coating of about the same age revealed 36 electrical leaks, 3 of which were examined. One defect was a large holiday beneath the wrapping; the second was a puncture of the coating by a ring from the pipe left in the trench; the third was a defective patch with the water beneath it. It will be noted that all of these failures were the results of poor inspection rather than of poor coating materials.

McCabe and Hull [153] reported that a test of 35,000 feet of 7-inch mastic-coated pipe showed an average resistance to ground of 500,000 ohms per linear foot about 6 months after it was placed in the ground. This may not have been a sufficiently long exposure to allow the coating to reach its minimum resistance.

As no one type of coating is superior to all others under all conditions, it is usually best to choose the type that can be most easily applied under the circumstances, because poor application is one of the chief causes of unsatisfactory performance.

## XIV. CATHODIC PROTECTION

Until comparatively recently, the methods discussed in the preceding sections were the ones commonly employed for the prevention or mitigation of underground corrosion. More recently, however, the method known as cathodic protection has been applied, and its use is growing rapidly in practice. Fundamentally, cathodic protection consists in impressing electromotive forces on an underground structure through auxiliary anodes in such a way as to make the entire structure cathodic with respect to the adjacent soil. Although this procedure does not eliminate

corrosion, it transfers the corrosion from the protected structure to the auxiliary anodes, which are more easily and economically replaceable. This method does not prevent direct chemical attack, which is ordinarily

negligible.

The use of zinc as an anode for the protection of ship bottoms, boilers, and similar structure has been practiced with varying degrees of success for many years. Also, a system of cathodic protection known as "pipe drainage" [19] has been used to protect pipe lines from corrosion due to stray currents from street railway systems. The beneficial effects were not fully appreciated, however, until the street railways were abandoned. In 1932, Scherer [154] reported that 26 companies were operating 542 cathodic protection units to protect 2,006 miles of pipe. Since that time, the number of miles of pipe so protected has considerably increased.

# 1. GENERAL PRINCIPLES

Theoretically, it should be possible to stop electrolytic corrosion by bringing the buried metal to such a potential with respect to the adjacent soil that no current flows out of the surface of the metal. Practically, however, neither the metal nor the soil can be made an equipotential surface and current will leave certain elements of the surface of the metal even though the average value of the current is zero. In order for all parts of the metal surface to be cathodic, therefore, the applied voltage must be great enough to cause a certain amount of current to flow into the protected metal surface. From the economic point of view, the current should be no greater than the minimum necessary to afford complete protection. Excessive current involves an unnecessary expenditure of energy and excessive consumption of anode material. The latter loss might be avoided by the use of noncorrodible anodes but this is generally not feasible.

#### 2. CRITERIA OF PROTECTION

The most difficult problem in connection with the application of cathodic protection is that of determining whether or not protection is actually effected. Obviously, the determination must be made indirectly because it is not possible to make direct observations of corrosion, and it would not be satisfactory to wait several years after the application of cathodic protection in order to determine by inspection whether or not it has been effective. Consequently, it has been necessary to adopt certain arbitrary empirical criteria, based for the most part upon experience, for judging whether or not the cathodic protection is adequate. The principal criteria in common use are based either upon estimates of the current density at the surface of the protected material or upon measurements of the potential of the protected material with respect to that of the soil. A third criterion that has been investigated is the relationship between the potential of the surface to be protected and the current flowing into it as the applied electromotive force is increased or decreased.

# (a) CURRENT DENSITY

Of the three criteria mentioned, current density at the protected surface is the least satisfactory for two reasons. (1) the value must be calculated as an average over a comparatively large area over which the distribution of the current is normally far from uniform, and (2) no simple way has been found for determining the value of the current density necessary in a given case. Furthermore, conditions are likely

to change with time, so that the value of the current density necessary

for protection may not be the same at different times.

The difficulty in determining the value of current density required for complete protection is apparent from the values given by different investigators. The values reported range from 1 to 15 milliamperes per square foot. Rhodes [155] found the minimum effective current density to be about 20 microamperes per square inch, or about 3 milliamperes per square foot, whereas Hill [156] from laboratory experiments concluded that the protective current density ranged from 1 to 10 milliamperes per square foot.

Rogers [157] reported that in one soil a current density of 3.65 milliamperes per square foot reduced the corrosion loss by 98 percent whereas in another soil a current density of 8.2 milliamperes per square foot reduced the loss only 82 percent. Keeling [158] found that a current density of 15 milliamperes per square foot was required to protect submarine pipe lines. Ewing [93] found that from 1.5 to 14 milliamperes per square foot was required to protect different sections of pipe in Louisiana and Texas.

A further complication arises if cathodic protection is to be applied to a pipe line on which there is a protective coating. Obviously, the current required would depend upon the number and size of the apertures in the coating. For instance, one operator found that the current required to protect a coated line was about 4 percent of that required for the protection of an uncoated line.

It appears from the experiences and opinions cited above that current density is not a safe criterion for determining whether or not cathodic

protection is adequate in a particular case.

#### (b) PIPE-TO-SOIL POTENTIAL

The purpose of cathodic protection is to prevent corrosion by making all points on the surface of the metal cathodic with respect to the adjacent soil. However, as the surface of the metal can not be made an equipotential surface, it is difficult to tell by any practical method whether or not the desired condition has been produced. On account of this uncertainty it is customary to make an allowance of from 0.2 to 0.3 volt to provide for inaccuracies of measurement and variations in service conditions.

The polarity of the metal with respect to the soil is usually determined by means of a high-resistance voltmeter, potentiometer-voltmeter or vacuum-tube voltmeter, and a steel or copper sulfate electrode. Some engineers follow the method of Rhodes [159] and use an electrode of pipe steel placed about ½ inch above the surface of the pipe. Values obtained by this method are said to be reproducible to about 0.02 volt. Sufficient current is applied to the pipe to make its potential 0.2 volt below that of the electrode. Other engineers [160-164] use a copper sulfate electrode placed above the pipe and assume that the normal potential difference between such an electrode and a pipe that is neither collecting nor discharging current is 0.54 volt [160]. They usually assume that the pipe is protected if its potential is lowered to —0.80 or —0.85 volt with respect to the copper sulfate electrode.

Records of leaks on one corroding pipe after cathodic protection has been applied and maintained for some time indicate quite definitely, at least under most soil conditions, that keeping the potential of the pipe line at least 0.85 volt below that of an adjacent copper sulfate electrode will prevent additional leaks after the protection has become established.

Many corrosion engineers feel, therefore, that this is a sufficiently good criterion for practical purposes, especially since the measurements can be made easily and quickly. However, if the pipe has a high-resistance coating with only small holes or cracks, this criterion may fail on account of the high IR drop involved in such cases.

There are several objections to the use of potential difference as a criterion of protection. Kuhn [165] has shown by an extended series of tests that the potential of cast iron with respect to a remote reference electrode varies from point to point. He also reported that the potential of the pipe as a whole changed with the time the pipe was in the earth. Gatty and Spooner [90] have shown that increasing the supply of oxygen raises the potential of iron in an electrolyte and that increasing the salkalmity of the solution lowers the potential of the iron. As soils differ widely in oxygen and alkali concentrations, they should also differ with respect to the soil-iron potentials.

That the potential of iron with respect to a copper sulfate reference electrode is not the same for all soils is indicated by the following tabulation. For a test of the effectiveness of zinc as a source of cathodic protection, three iron rings were buried in each of eight locations. The potential of each ring was measured with a potentiometer-voltmeter before the experiment was started. The potentials at six of the test sites are shown below:

Location	Potential of iron ring with respect to CuSO ₄ electrode Volts
Austintown, Ohio	0.575 to _ 0.600
Deerfield, Óhio	
Rocky Ford, Colo	
Albuquerqué, N. M	
Los Ángeles, Calif	
Shreveport, La	—.753 to —.760

These differences in potential may be accounted for in part by different percentages of anodic areas, as they are the resultants of the potentials of the anodes and cathodes on the surface of the iron.

The potential difference between a pipe and an electrode some distance from its surface may be affected by an IR drop caused by current flowing to or from the pipe or by other currents in the earth. This may be important in cases of cathodic protection. On account of the possible sources of error just mentioned, many engineers use the change in potential of the pipe when current is applied as the criterion for the degree of protection and assume that the pipe is protected when its potential has been lowered 0.3 volt. Obviously, the condition of the pipe may be such that no current is required for its protection, in which case the lowering of its potential is unnecessary. Under other conditions the change in potential of the pipe with respect to an electrode some distance away may be due largely to the IR drop in the soil and not representative of the change at the pipe surface. To avoid this error some engineers use an electrode separated from the pipe by only about 1/8 inch of insulation. This avoids most of the IR drop unless the pipe has an insulating coating, but such measurements are influenced by the condition of the pipe adjoining the electrode. It is, of course, possible to take account of the resistance of the soil or coating surrounding the pipe, as has been done by Ewing [164] and Scott [166, 167]. The most important objection to potential or change in potential as a criterion for protection was pointed out by Ewing [93], who showed that when current was applied to a cathode such as a pipe line, the potential of the protected metal with

respect to a reference electrode might gradually increase or decrease, depending on the reaction at the cathode. The reaction is determined by the aeration and pH value of the soil.

# (c) CURRENT-POTENTIAL CURVES

In the course of a study of the velocity of corrosion Britton [92] found that when the current flowing to a piece of iron in aerated 1/10 N potassium chloride solution was gradually reduced from a value more than sufficient to prevent corrosion of the iron, the potential of the iron with respect to a reference electrode quite near it increased up to a certain value and then remained constant as the current was further reduced. Experiments as well as theoretical considerations showed that the break in the curve occurred at the current density just sufficient to prevent corrosion. The experiment was repeated with other concentrations of potassium sulfate. The explanation of the horizontal part of the curves, where there was no change in potential of the cathode with increasing current, is that the truly cathodic areas were large compared with the corroding areas and so determined the measured potential. This potential did not change because all the current entered the anodic areas. It was used in counteracting the local or corrosion current from the anodes. This unequal distribution of current may be explained by the assumption that the cathodic area was insulated by a relatively high resistance film.

If the corrosion is not controlled by the cathode, the current potential curve may not have a horizontal part or even a critical point. However, the corrosion of iron is usually controlled by polarization of the cathodic areas.

Ewing [93], working at the National Bureau of Standards as a Research Associate for the American Gas Association, undertook an extended investigation to determine how the phenomenon discovered by Britton could be used to determine the current required for the protection of pipe lines. Müller [89] showed that the measured potential of a corroding cathode was its true potential diminished by the IR drops resulting from the flow of the local and imposed currents, i.e., that if these IR drops were deducted or eliminated from the observed values, the true value would be obtained. Ewing also showed by reasoning and citations of the work of other experimenters [168] that after the current became sufficient to prevent corrosion, the cathode would act as a hydrogen electrode and that the potential of such an electrode varied linearly with the logarithm of the current flowing to the cathode. Consequently, if the true potential of the cathode is plotted against the logarithm of the impressed current, the resultant curve should consist of horizontal and sloping lines connected by a curve representing the transition to the overvoltage curve, as shown in figure 77. The intersection of the projections of the straight lines is taken to indicate the current required to prevent corrosion. If the resistance of the electrolyte is low, sharp breaks in the current-potential curves can be obtained without correcting for the IR drop, as was done by Britton. With high-resistance electrolytes, however, a smooth curve with no break is obtained (see fig. 79) if the potential is not corrected for IR drop. A laboratory method for obtaining data for current potential curves free from IR drop has been described by Pearson [169]. His apparatus was simplified and made applicable to field work by Hadley [170] and associates. Figure 78 shows the essential features of the Pearson-Hadley circuit. At the top of the figure is a source of variable current, consisting of a battery and resistance connected through an ammeter and rheostat between an anode and the pipe to be protected. The remainder of the circuit consists of two copper sulfate electrodes, E+ and E-, a high-resistance null indicator, V, preferably a vacuum-tube voltmeter, a potentiometer-rheostat,  $R_3$ , and a potentiometer voltmeter, referred to later as a back-out circuit. The operation of the apparatus is as follows: With the battery circuit open and switch  $S_2$  in the down position, the potential of the pipe with respect to electrode E- is obtained by applying a counter-potential by the backing-out apparatus  $R_2$ , until indicator V reads zero. (If V is a calibrated vacuum-tube voltmeter, it will indicate the potential if the backing-out potential is set at zero.)  $S_2$  is then reversed, the current cir-

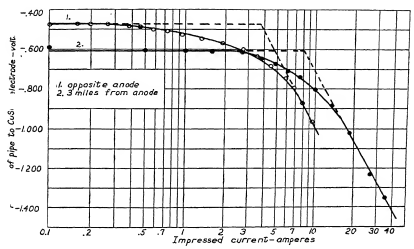


FIGURE 77 .- Current-potential curves for 6-inch pipe.

cuit closed, and rheostat  $R_1$  adjusted to permit the passage of the estimated protective current or any other current sufficient to cause a well-defined change in the reading of the vacuum-tube voltmeter when switch  $S_1$  is shifted. The switch is then moved from side to side, thus changing the current, and the sliding contact, N, on the null potentiometer,  $R_3$ , is adjusted until changing the position of switch  $S_1$  causes little or no change in the reading of the vacuum-tube voltmeter. When a change in current does not affect the deflection of the voltmeter, the IR drops in the earth are just balanced by the corresponding IR drop in the null potentiometer, and their effect is eliminated from the result.

A comparison of the null circuit, 1, with the Wheatstone bridge circuit, 2, and the equivalent null bridge circuit, 3, in figure 78, in which corresponding points have been similarly lettered, may make the operation of the circuit clearer. The batteries in the null bridge circuit represent the voltage supplied by the backing-out potentiometer and the potential of the pipe with respect to the reference electrode. If the bridge is first balanced with respect to resistances (IR drops), the galvanometer, V, will read zero when the backing-out voltage equals the potential of the pipe. It is advisable to make the positions of E— and E+ such that

a balance is obtained when the contact, N, is somewhere near the center of  $E_3$  since the apparatus is most sensitive to changes in I under this condition.

When the bridge or null circuit has been balanced for a current high enough to give a reasonable sensitivity, the current is reduced to a very small value. After sufficient time has elapsed for the vacuum-tube voltmeter to reach a steady state, the potential across it is backed out by

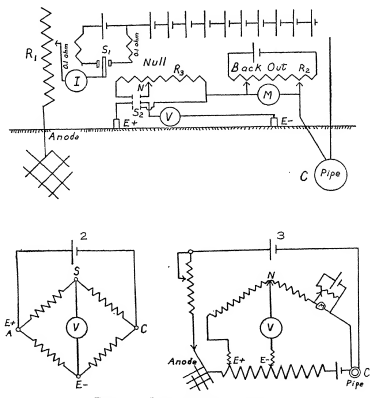


FIGURE 78.—Pearson-Hadley null current.

1, Null circuit; 2, wheatstone bridge current for comparison with 3, the testing circuit in 1. The letters designate corresponding parts in each circuit.

means of the potentiometer resistor,  $R_2$ , and the readings of the ammeter, I, and the voltmeter, M, are recorded. Rheostat  $R_1$  is then readjusted to pass from 30 to 50 percent more current and a new balance is obtained. Thus a series of observations of potential and current are accumulated. At first, little or no change of potential will be observed. Observations should be continued until four or more definite changes in potential have been observed.

Usually, it will not be necessary to readjust the null circuit until after the critical current has been passed. In some cases, the circuit becomes badly unbalanced for large currents. It is advisable, however, to check the balance of the null circuit from time to time. The data for figure 77 were obtained by Hadley on a 6-inch pipe. The apparatus used in obtaining the data for curve 1 was set up near the point where the power was connected to the pipe, but the anode was about a quarter of a mile from the pipe and the positive copper sulfate electrode about 125 feet from the pipe. A welding generator supplied the current.

The voltage data for curve 2, figure 77, are for a point on the same pipe about 3 miles from the power connection. The welding generator, ammeter, and control rheostat were at the same place in each test. In the second test, communication was by radio, but a modification of Smith's communication system described in Ewing's book [29], under the topic "Measurement of Network Constants," could have been used. The negative copper sulfate electrode was placed over the pipe and the positive electrode was about 90 feet from the pipe. Curve 2 is presented to show that the null method can be used to obtain data at points remote from the power connection.

The potential-log current curve consists of but two straight lines only when little or no current flows to the cathodic areas on the pipe while anodic areas remain. This condition obtains when the corrosion is cathodically controlled, i.e., where corrosion products on the cathode determine the rate of corrosion either by their resistance or potential. If cathodic control is incomplete, the curve for low current densities will not be horizontal but have a slope. If the cathodic control fails gradually, the potential-log current curve will bend. Bending will also occur if the potential readings contain an IR drop. For this reason, it is important to maintain the balance of the null circuit.

That part of the polarization potential that is the result of the accumulation of alkali at the cathode changes gradually and persists for a considerable period after the current has been cut off. Because of this type of polarization, observations of current and potential must be made at equally spaced time intervals if a smooth curve is to be obtained. Two or three minutes is a good interval, which allows sufficient time for the adjusting of apparatus and occasional checks of the null circuit. This persistence of polarization accounts for the fact that the operator cannot drop back to a lower current to check an observation in the course of a test and also for the fact that the potential at zero current after the test will not be the same as initially. Successive tests within a few hours of each other will often show small decreases in the protective current for the same reason.

Tests run with the initial current either at the maximum or at the minimum value will give substantially the same value for the protective current, but the potential for zero current after a test will be lower than the original potential of the pipe. For this reason the potential of the pipe should be measured before any current is applied.

The laboratory tests upon which the above method of determining the protective current is based were performed with small electrodes in a uniform electrolyte. It might be asked whether or not they are applicable to pipe line along which the the potential varies. Pearson [171] has pointed out that 90 percent of the observed potential of a line with reference to an electrode at the surface of the ground is contributed by a segment four times as long as the distance from the pipe to the electrode.

An electrode placed over a line 30 inches deep, therefore, is affected by the potential of about 10 feet of the line and is only slightly affected by the remainder of the line. To determine the current required to protect this part of the line, it is necessary to measure the current in the pipe on either side of the segment and so compute the current collected by the segment under observation. The fact that, with care and adequate apparatus, log current-potential curves with reasonably sharp critical points can be obtained indicates that the laboratory test is applicable to field conditions.

Under some conditions, the potential-log current curve consists of three or more parts. The work of Gatty and Spooner [90] suggests that if the cathode has been exposed to oxygen, the first change in the slope of the curve represents a reduction of the oxides on the cathode. When this reduction is complete, the hydrogen-overvoltage curve follows. If the current density is increased sufficiently, the curve may again depart from its previous course.

Some unpublished work by Denison suggests that the true protective current is indicated by the point where the potential curve first begins to bend rather than by the intersection of the horizontal and the hydrogen over-voltage lines, and this is in line with Miller's unpublished opinion that the latter point sometimes indicates a current greater than that necessary for protection.

# (d) COMPARISON OF RESULTS OF DIFFERENT METHODS FOR DETERMINING THE PROTECTIVE CURRENT

As the apparatus required for the Pearson-Hadley method for determining the current required for the protection of a buried structure is obviously more complicated than that in common use, and since the time required for the tests is longer, it is of interest to consider whether or not the different methods give similar results, as well as to consider the evidence for their validity.

Figure 79 shows current-potential curves taken in several ways. The cathode was a buried tank with 8- by 12-inch steel places placed about 8 inches from the tank near the top, middle and bottom. The sides of the tank were coated with coal-tar enamel.

Curves 1, 2, and 3 show the potential of the tank with relation to the steel reference electrodes near the top, center and bottom of the tank, respectively. It will be noted that the tank is at nearly the same potential with repect to the top and bottom electrodes, and that there is a difference of potential of nearly 0.3 volt between the top and middle electrodes.

If the tank is protected when its potential is 0.3 volt below that of the reference electrode, 1.3 amperes are required for protection, according to the indication of the central electrode, while the other electrodes indicate that no current is required. If a change in potential of 0.3 volt with respect to a steel electrode is the criterion, the required currents are 1.4, 1.5, and 1.9 amperes for the top, center, and bottom electrodes, respectively.

If a copper sulfate electrode is the reference point, curve 4 indicates that 0.32 ampere is the protective current on the basis of a potential difference of 0.85 volt and 1.3 amperes on the basis of lowering the potential 0.3 volt. If the null method is used, curve 6 indicates that a current of 0.7 ampere is required for protection. Other curves obtained by the null method indicated that the protective current was 1.0, 1.1, and 1.2 amperes.

For the tank under discussion it appears that the change of 0.3 volt and the null method give about the same results within the limits of reproduction. Changing soil and electrode conditions made it impossible to determine a single value of current which will just furnish protection at all times so far as the tank in question is concerned, and prob-

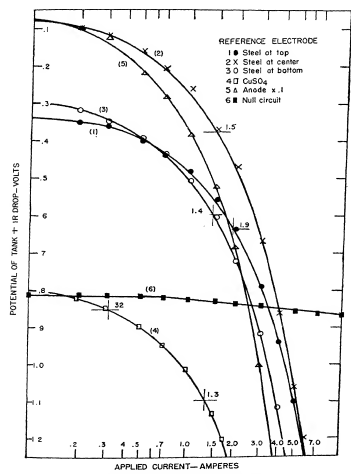


FIGURE 79.—Current-potential curves for a buried tank.

ably an engineer would have chosen about the same value for the protective current for the tank whatever his criterion.

Similar conclusions do not always result from such comparisons of methods. Figure 80 shows the results of two tests run simultaneously by Hadley and others on a 40-foot section of bare 16-inch steel pipe that was separated from the remainder of the line by Dresser coupling. The

upper curve was obtained by the null method. It indicates that 3 amperes are required to protect the section. A similar test run a few hours earlier indicated 2.5 amperes were required for protection. The lower curve was obtained by the usual method, using a potentiometer voltmeter and a steel electrode placed  $\frac{1}{16}$  inch above the pipe and a foot to one side of the copper sulfate electrode used for the null method. The curve indicates that 18 amperes are required to lower the potential of the pipe 0.3 volt. A third simultaneous test employing another steel electrode 1 foot on the other side of the CuSO₄ electrode indicated that 30 amperes were required to lower the potential of the pipe 0.3 volt. A protective system installed by a consulting engineer was furnishing 0.1 ampere for the protection of this section.

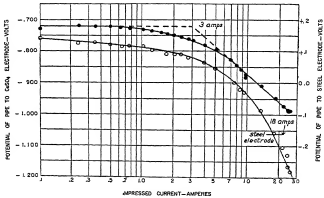


FIGURE 80 .- Current-potential curves for 40 feet of 16-inch pipe.

When a pipe line is made cathodic, two changes occur at or near its surface. (1) There is polarization resulting usually from the deposition of hydrogen. Most of this change occurs within a few minutes and disappears within a short time after the current is cut off. (2) There is an accumulation of alkali near the surface of the pipe. This change increases with the flow of current until it is neutralized by the diffusion of the alkali into remote soil. The effect of the alkali persists to a decreasing extent for some time after the current is cut off. Because of these changes it is not possible to make the most economical adjustment of the protective current until the system has been in operation for some time. The current supplied by the source of power will be affected by the everchanging resistances of the anode, cathode, and intervening soil, and a redetermination of the current required for protection should be made from time to time if the use of a minimum current is desired.

Obviously, the amount of current needed will depend on the exposed area of the pipe. For this reason, any insulating coating on the pipe tends to reduce the current necessary to complete the protection.

Complete protection depends on making every point on the surface of the pipe cathodic, that is, on making current flow to all points. Those familiar with the distribution of current over the surface of an electrode know that it is difficult to force current into or out of a narrow crack or depression. It follows that it will require more current to protect a badly pitted pipe than one with a smooth surface. It is very difficult to protect a deeply pitted pipe. This accounts for the fact that many users of cathodic protection find that leaks do not stop for several months or years after protection is applied to a badly corroded line. This is illustrated in table 100, which shows the number of leaks that developed in protected section of pipe lines totaling approximately 36 miles [172]. The criterion for protection was a potential of 0.8 volt or more between the pipe and a copper sulfate electrode on the surface of the ground over the pipe.

	•
Exposure period after application of cathodic protection	Number of leaks
Years 1 2 3 4 5 6	130 242 17 8 7 6

Table 100 .- Effect of cathodic protection on rate of leak occurrence.

Some of the leaks during the 2 years following the application of cathodic protection may be attributed to imperfect adjustment of the protection apparatus, and it may be that more current is required initially than later, but it seems probable that the continuous occurrence of leaks after protection was applied may be due to the fact that the protective current could not be forced into the bottom of the deep pits. When these were eliminated by their development into holes, which had to be repaired, the current was sufficient to protect the pipe.

# 3. CATHODIC PROTECTION INSTALLATIONS (a) PROTECTION OF TRANSMISSION PIPE LINES

(1) Principles.—Fundamentally all that is required for cathodic protection is a source of direct current and a means of getting the current into the earth and off of the pipe to be protected. Cathodic protection engineering consists in determining how much current is required and selecting the most economical means for supplying and distributing the current. In general, two systems of current distribution are used, regardless of the source of the current. One system introduces the current into the earth at a single favorable location, and attempts thereby to protect as much pipe as is practicable. The current density at the surface of the pipe and the potential of the earth with respect to the pipe is greatest near the point where the current is removed from the pipe, which is usually the point on the pipe nearest the anode. The distribution of the current flowing to the pipe tends to become more nearly uniform and the amount of protected pipe tends to increase with increased separation between anode and pipe up to distances of a few hundred feet. This system requires the minimum wire to transmit current but is wasteful of current in the region near the anode. The distributed anode system, patented by Rhodes [155, 159, 173], requires less power but a greater investment in copper and power converters. For zinc as a source of current, the Rhodes system is almost essential, and for city networks it has certain advantages.

The design of cathodic protection for cross-country lines has been discussed by Rhodes [155], McGary [174], Rogers [175], Thayer [176],

Scott [167], Ewing [164], Schneider [177], and Olson [178]. Three main points have to be considered (1) an economical source of power including distribution costs and losses (2) a low-resistance anode, preferably with little polarization, low counter voltage or over-voltage and long life (3) as extensive a distribution of current from a single source as is consistent with economy, and with no interference with unprotected structures. Some of these requirements will be discussed in detail.

(2) Sources of Current.—(a) Converted Power. Windmills, gas-engine-driven generators, and rectifiers have been in use so long that their design has become stabilized, and to obtain satisfactory apparatus of these types it is only necessary to select a reliable manufacturer and furnish him information as to the current and voltage which are required.

Scherer estimated that in 1941 there were 750 cathodic units protecting 3,000 miles of pipe line. Approximately 45 percent of the total number of installations used rectifiers with outputs from 50 watts at 6 volts to 1 kilowatt at 20 volts. Their popularity is due in part to the fact that usually they contain no moving parts and require little attention other than an occasional adjustment of voltage. Single-phase copper oxide rectifiers have an efficiency of about 65 percent [179] and three-phase apparatus an efficiency of about 85 percent. Selenium and copper sulfide rectifiers have about the same efficiency. Electron-tube rectifiers are reported to be somewhat more efficient.

Gas-engine-driven generators are used frequently to protect gas lines, where gas to operate them can be taken from the line. Wind-driven generators furnish power for about 20 percent of the installations. Their popularity seems to be decreasing largely, because they have been installed where the wind is sufficiently continuous to be a satisfactory source of power. The velocity of the wind should not fall below 5 miles per hour for periods longer than a day, and the windmill should operate at least 75 percent of the time. The electrical and mechanical apparatus requires little attention. United States Weather Bureau reports are of value in a preliminary survey of wind possibilities in a given region but sometimes they are not applicable to the spot where a windmill is desired.

(b) PRIMARY POWER.—In sections of the country remote from power lines and unfavorable to wind-generated power, and in cities where the stringing of wires to supply current would be unsightly or expensive, and where anodes must be placed close to the protected structure to avoid endangering neighboring structures, the anodes themselves may be made to furnish current if the resistance of the soil is sufficiently low. Aluminum, magnesium, and zinc have been suggested because their potentials are less noble than steel.

A. ALUMINUM ANODES.—The electrode-potential series indicates that aluminum might be more effective than zinc as a current-generating anode for cathodic protection. Aliter [180] has reported an experiment in which he compared the effectiveness of the two metals for the protection of steel rods in two Hanford soils. His results indicated that the aluminum was more effective in a soil containing carbonates in which the zinc reversed its potential with respect to iron. Both the zinc and the aluminum contained iron as an impurity. As aluminum is attacked by some alkali soils, its use in such soils might be limited by local corrosion, although the pH of the Hanford soils was from 7.7 to 8.1.

B. MAGNESIUM ANODES.—The open-circuit potential between iron and magnesium is in the order of 1 volt, which is more than twice that

between zinc and iron. Olson [181] has reported some experimental work on magnesium anodes. Some of the anodes became inactive after a few months of exposure. Others continued to supply 50 percent or more of the original current. However, the loss of weight of the magnesium anodes was several times that attributable to the current supplied to the cathode (pipe). This investigation is being continued in an effort to find a magnesium anode less subject to local or galvanic corrosion. Grebe and McNulty [182] have reported extensive experiments with magnesium anodes, which show electrolytic corrosion efficiencies as high as 30 percent. They predict that future developments will make the use of magnesium anodes economically sound.

c. ZINC ANODES.—The open-circuit potential difference between iron and zinc is in the order of 0.4 volt, which may be enough to furnish an adequate protective current if the resistances of the soil and anode are low. Zinc anodes have been installed in a number of localities with beneficial effects, though in some cases the protection which they have furnished has not been complete. These failures to protect are in part attributable to an insufficient number of anodes and in part to improper installation or maintenance. Reports on zinc anodes have been published by Rhodes [159], Smith and Marshall [183], Brockschmidt [184] and Mudd [185]. Much of the work of these men and considerable additional information on zinc anodes has been summarized in a report by Wahlquist [186], from which much of the following paragraph was abstracted.

Zinc anodes were attached to a gas line in eastern Colorado in 1935. The first rods were 1 inch square and 4 feet long with a ¼-inch iron core, and each weighed 12 pounds. Later rods were round, with a diameter of 13% inches, and each weighed 18 pounds. The rods were cast locally from electrolytic zinc. Usually eight rods, spaced about 10



FIGURE 81.—Zinc carbonate formed from a zinc anode used to protect a pipe line.

The diameter is 1½ inch.

feet part, were connected together in a line parallel to the pipe line and from 5 to 14 feet from it. Most of the soils contained sulfates and carbonates and had resistivities between 700 and 200 ohm centimeters. Most of the installation furnished between 30 and 50 milliamperes per rod after from 6 to 8 years and maintained the pipe-to-soil potential at from -0.6 to -0.93 volt with respect to a remote copper sulfate electrode. Several leaks have occurred since the installation of the zinc anodes, but the frequency of their occurrence has been greatly reduced. The condition of the rods indicates that they will continue to furnish current until the zinc has been nearly consumed or until the iron core is exposed. The most common corrosion product is zinc carbonate, which remains in place around the rod as a moist solid cylinder. Figure 81 shows a section of such a carbonate cylinder pulled from an old rod. The wall thickness of the carbonate is  $\frac{36}{2}$  inch.

As a result of his experience, Mudd [185] uses a zinc anode rolled

from 14-inch sheet zinc to form a cylinder 7 inches in diameter and 3 feet long. He places these cylinders in holes 8 inches in diameter and from 4 to 6 feet deep. Calcium-sulfate (gypsum) is placed around the cylinder, which is packed with crushed rock to expand the cylinder and make a good contact with the earth. Mudd found that his anodes were most effective when several were connected in parallel to give a total anode area of 30 square feet. He placed his nearest anode more than 100 anode diameters from the pipe and made the spacing between anodes 50 diameters, conforming to Schneider's [187] recommendations. Mudd's experience is not quite so favorable to zinc anodes as is that of Wahlquist [186].

Brockschmidt [184] connected ten 4 foot by 1%-inch zinc anodes in parallel at 5 foot intervals in a line 3 to 5 feet from an 18-inch Dresser-coupled enamel-coated gas line in the first bottom of the Mississippi River. Most of the soil had a resistivity in the neighborhood of 500 ohm-centimeter. After about 2½ years the average potential of the pipe was -0.198 volt with respect to a steel electrode in the soil a fraction of an inch distant, and -0.990 volt with respect to a remote copper sulfate electrode. After 449 days the total output of 10 electrodes was 110 milliamperes. The pipe was still positive to earth at one location.

The output of Brockschmidt's rods was less than half that of those in the Colorado installation, where the size and location of the rods and also the soil resistivities were approximately the same. Since the pipe-to-soil potentials were generally greater than the commonly accepted criterion of -0.85 volt for safety, the two installations may have been equally effective. Only the leak records will determine whether or not this is the case. The data presented as to the effectiveness of the two installations are unsatisfactory because they show only averages of widely variant data and because the criterion for effectiveness has not been established. There can be little doubt, however, that under many conditions zine anodes properly installed and maintained will materially reduce corrosion losses.

Rhodes [159] and some others have stressed the importance of the use of very pure zinc for anodes. Others have not found this necessary. Additional research and experience will be necessary to determine the limitations of zinc anodes. Among the limiting factors are (1) low potentials, which limit their application to pipes in low-resistance soil, (2) local or galvanic corrosion, (3) the occasional development of high-resistance corrosion products, and (4) the reversal of potential (confined probably to impure zinc). As has been said, zinc anodes are almost necessarily placed close to the pipe, and as a result they have the advantage in city streets of not endangering adjacent unprotected structures. Another factor in their favor is the comparatively uniform pipe-to-soil potential resulting from the use of the distributed anode system.

(3) Anodes (a) RESISTANCE TO EARTH.—In the preceding section, anodes have been discussed only as sources of potential. In the following section they will be treated as means of getting current into the earth. As such, their important characteristics are (1) resistance to the flow of current, i.e., electric resistance to earth, polarization, over-voltage, and counter electromotive force, (2) permanence, and (3) cost.

The resistance of the anode itself is usually negligible in comparison with that of the remainder of the electric circuit. Most of the resistance to the flow of the protective current is in the earth at or near the anode and cathode. If the cathode, the pipe to be protected, is bare, the resis-

tance at its surface is usually small and attributable to corrosion products or polarization. Making the resistance between anode and cathode low is one of the major problems of cathodic protection. This depends

largely on the size, shape, and location of the anode.

Frequently, several anodes are connected together in a line parallel to the pipe to be protected, in which case they are known as distributed anodes, or in a two-dimensional group known as a ground bed. The distributed anodes usually consist of rods from 1 to 2 inches in diameter and 4 to 10 feet long. Eight or more are spaced from 5 to 10 feet apart in a line parallel to and a few feet from the pipe. The resistance to earth of the group is approximately that of the resistances to earth of the individual anodes in parallel, that is,  $1/R = 1/r_1 + 1/r_2 + \dots 1/r_n$ , where R is the resultant resistance and  $r_1, r_2$ , etc. are the individual resistances to earth. The same equation applies to any other arrangement of anodes if they are sufficiently separated from each other.

The resistance to earth of the individual anodes depends on, but is not proportional to, their diameters and lengths and on the resistivity of the soils in which they are placed. The anodes may be placed vertically or horizontally, the choice depending largely on the depth of relatively low resistance soil. The data on the resistance to earth of ground rods for electric systems are applicable to anodes if they are placed in homogeneous soil. This subject has been discussed by Peters [188], but the additional resistance resulting from heating, osmosis, and polarization

must be added.

Peters [188] showed that the resistance to earth of a pipe driven in the earth was represented by the equation  $R = \varrho/2\pi C$ , in which R is the resistance between the pipe and the earth,  $\varrho$  is the specific resistance of the soil, and C is the combined electrolitic capacity in free space of the electrode and its image above the surface of the ground. The value of C can be calculated approximately from the equation C = 2L/2 log. (4L/d), in which L is the length of the pipe and d its diameter. The equation can be used for computing the effect of increasing the diameter or length of the pipe. Figure 82 illustrates the effect of increasing the length of the pipe as determined by the above equation and by experiment in a high resistance soil.

Frequently, the resistance of the subsoil or underlying rock is so high that extending a ground rod into it does little good. It is advisable, therefore, to determine the resistivity of the soil at various depths by Gish and Rooney's [189] application of the Wenner method, as described

earlier, and to limit the length of the anodes accordingly.

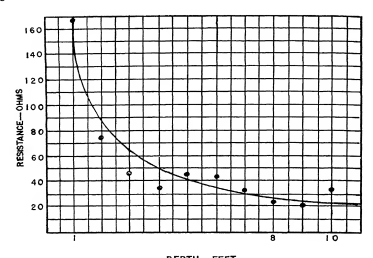
The effect of putting two or more vertical rods in parallel is illustrated in figures 83 and 84. The effect of the area of a plate near the surface

of the earth on its resistance to earth is shown in figure 85.

If the subsoil has a high resistance, it may be advisable to lay the anode horizontally, in which case its resistance to ground will be  $R = \varrho \log_e(2L/d)/\pi L$ , in which the letters have the same significance as in

the preceding equation.

If the resistivity of the soil is constant, the resistance to earth of a horizontal ground plate, such as a sheet of metal, decreases as its distance below the surface of the earth increases until, at depths large in comparison with the diameter of the plate, it is half that at the surface of the ground. The resistance to earth of a circular plate at the surface of the ground is R=r/2d, in which r is the resistivity of the soil and d is the diameter of the plate. Consequently, to reduce the resistance to earth of a plate to one-half, its area must be made four times as great. It would



DEPTH-FEET

FIGURE 82.—Effect of the length of a driven-pipe ground on its resistance to earth.

be more economical to use two small plates in parallel and some distance apart. Likewise, a number of small widely separated rods or strips in parallel have a lower resistance to earth than a continuous piece having the same area.

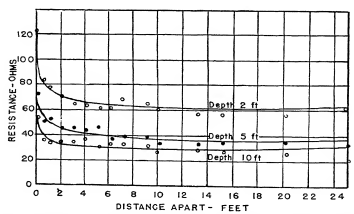
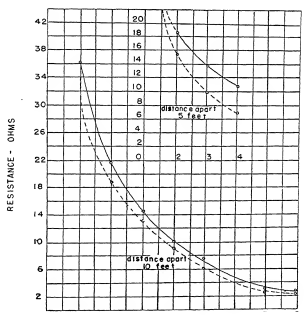


FIGURE 83.—Effect of the distance apart of two driven-pipe grounds in parallel on their resistance to earth.

(b) Materials. A. Iron.—Because of its low first cost, low over-voltage and the low resistance of the corrosion products, junk iron, usually old pipe, is often used for anodes. When this is done, care should be taken to separate the individual pieces as much as possible and so to interconnect them that a break in the continuity of the group will not make

part of the electrodes useless. More than one connection to each piece should be made.

The kind of iron or steel used is unimportant. Boilers and parts of old engines and large machines have been used and also old well casings in place. If the current density at the surface of the iron is kept low, the anode will lose about 20 pounds per ampere-year plus some loss caused by local galvanic action. The latter will depend on the corrosivity of the soil and the homogeneity of the metal. If the current density is high, the rate of loss of metal may be less, since part of the current may cause the evolution of oxygen by the electrolysis of water in the soil. Additional energy will be required to ionize the water and still more to overcome



NUMBER IN PARALLEL
FIGURE 84.—Effect of the number of 10-foot driven-pipe grounds in parallel on their resistance to earth.

Observed values

— — — — = Observed values. — — — O — — — = Calculated values.

the resistance resulting from the heating and drying of the soil. Usually, the cost of the electric energy will exceed the cost of the metal saved and on this account it is generally advisable to keep the current density low

The relation of current density to corrosion efficiency has been investigated by McCollum and Logan [190], who found that the ratio of the actual to the theoretical loss of anode material ranged from 1.40 for a current density of 0.1 square centimeter to about 0.4 for a current density 20 times as great. They found also, that the addition of acids, alkali, and salts to the soils, materials which affect the self-corrosion of

iron, had little effect on the efficiency of electroyltic corrosion. An exception to this is the effect of the addition of chromates. This greatly reduces the efficiency of electrolytic corrosion, probably because the iron becomes passive. They found no definite relation between the electrolytic corrosion efficiency and the rate of self-corrosion. This indicates that the factors affecting the Corfield [54] corrosivity test are chiefly soil resistivity and polarization. The low corrosion efficiency when chromates are present suggests a means for making a substantially noncorrodible anode.

B. CARBON AND GRAPHITE.—When iron is not available, carbon or graphite rods are frequently used. Rhodes [159] and Roddey and Shepard [191] at one time used arc-furnace electrodes 2 inches in diameter and 6½ feet long. They were placed in holes 6 inches in diameter and 10 feet deep and surrounded by crushed coke or coke breeze. The resistance of these grounds ranged between 1 and 4 ohms.

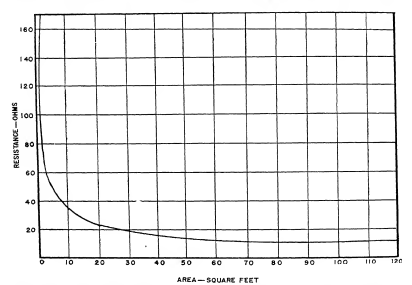


FIGURE 85.—Effect of the diameter of a buried circular plate on its resistance to ground.

To the IR drop caused by this resistance must be added a counter voltage of about 0.8 to 1.6 volts because of the potential of carbon with respect to steel.

According to Roddey and Shepard [191], the conduction between the carbon rod and the crushed coke is metallic and the rod does not deteriorate. There is a loss of the coke. A test which they ran indicated a loss of 0.000164 gram of coke per coulomb or 11.4 pounds per ampereyear. They attribute this loss partly to the formation of carbon dioxide, but mostly to the migration of the coke into the soil. The rate of loss of carbon anodes given above is greatly in excess of the value of 1.5 pounds per ampere-year given by Dorcas [192]. The latter is a

theoretical value based on the assumption that the loss of carbon is due entirely to the formation of carbon dioxide.

c. NONCORRODIBLE MATERIALS.—The cost of anode materials and their replacement has led to several suggestions for noncorroding anodes [193].

The user must decide whether it is cheaper to pay for material that must be replaced or for power required to generate and discharge oxygen from a noncorrodible anode. In certain cases an insoluble anode may be desired in order to avoid contamination of the electrolyte by corrosion products. Stainless steel and high-silicon cast iron have been suggested for this purpose, but few data applicable to cathodic protection are available.

# (b) CATHODIC PROTECTION OF CITY NETWORKS

Most cathodic protection has been applied to cross-country lines under conditions that make interference with other lines improbable or easy to prevent. When one pipe system in the same street with other underground pipe or cable system is protected by causing current to flow to it, this current may flow onto and off the adjacent structures and damage them. The adjacent structures may also interfere with the desired distribution of current. It is necessary, therefore, to take special precautions if cathodic protection is applied to city networks.

Interference has been discussed by Smith [194]. The application of cathodic protection to city networks has been described by Kuhn [195], Schneider [196], and Wainwright [197]. It is advisable that all pipe owners in the neighborhood of a proposed cathodic protection be informed of its installation and be given an opportunity to participate in the work or at least to determine whether or not it affects their properties. A tentative plan for such cooperation has been proposed [198].

Two solutions to the problem of cathodic protection in cities may be considered. Often the preferable one is the protection of all structures in a neighborhood by the joint action of all interested parties. This involves an agreement as to the distribution of costs and adjustment of potentials, which is occasionally difficult to determine satisfactorily. The alternative is the application of protection to a single structure. If this is done, a current and potential survey or an exposure survey of all structures that may be affected should be made both before and after cathodic protection is applied. Anode locations and bonds or insulating joints should be adjusted until no structure is endangered. In some cases the least interference will be caused by a system of distributed anodes placed close to the pipe to be protected. This system has been patented by Rhodes [199], who permits its use without royalties under very reasonable conditions. However, arrangements for its use should be made in advance.

# (c) CATHODIC PROTECTION OF TANK BOTTOMS

The protection of the bottoms of oil- and gas-storage tanks and the attached pipe systems is not different in principle from the protection of pipe lines. The same criteria for protection are used. Rectifiers are the most common source of power.

The protection of tanks has been described in some detail by Holsteyn [162], Brannon [163], and Bond [200]. The designs of the protection were based largely on the results of preliminary experiments sup-

plemented by experience in protecting pipe lines. Any one having a similar problem will receive considerable help from a careful reading of these papers.

#### (d) COST OF CATHODIC PROTECTION

The cost of cathodic protection will, of course, vary greatly with local conditions. A first approximation of the costs may be based on the experiences of others, but the cost of any installation and the effectiveness of the protection will depend to a large extent on the skill, experience, and foresight of the engineer in charge of the installation. For this reason, it is frequently economical to employ a consulting engineer to

design and install the first system.

Wainwright [197] estimates the installation costs for an isolated anode system for a city distribution network as \$700 to \$1200 and the annual charges as 29 percent of the installation cost. For an installation of zinc anodes in Colorado, estimated to have a life of 20 years, Rhodes [199] reported a cost, including development charges, of \$340 per mile plus \$2.62 per anode. He used from 240 anodes per mile for a 4-inch pipe to 680 anodes per mile for 12-inch pipe. Wahlquist's [186] examinations indicated that in some soils 20 years is too long an estimate for the life of anodes of the size Rhodes used. The amount of power required for protection has already been discussed. The amount depends on the condition of the coating, the criterion for protection, and the position of the anode with respect to the pipe.

Roddey and Shepard [201] reported current requirements of 13.1 amperes per mile of 22-inch enamel-coated pipe and 8.3 amperes per mile of 14-inch felt-shielded asphalt-coated pipe. They report an average earth-to-pipe potential of about 0.3 volt; position and kind of reference electrode were not stated. Olson [178] reported that it required from 4.25 milliamperes per square foot to 1.5 milliamperes per square foot to protect bare gas lines in Louisiana and Texas. His criterion for protection was not stated, but his published curves indicate that he endeavored to make his pipe negative to a copper sulfate electrode by at least 0.85 volt.

Schneider [202] has prepared a digest of cathodic protection equipment costs. The economics of cathodic protections have been discussed by Thayer [203], who describes several sources of power and concludes that for every dollar spent for protection his company has saved from eight to ten dollars in replacements.

For the protection of a bare 10-inch line, Stewart [204] reports the use of thirty-four 8-volt 24 ampere rectifiers spaced a third of a mile apart. The annual cost of power was \$0.013 per foot per year. On another bare line that was maintained 1-volt negative to a copper sulfate electrode, the current density ranged between 1.5 and 5 milliamperes per square foot. There were no leaks. Stewart's practice is to create a pipe-soil potential of at least 0.3 volt in excess of the potential existing before protection was installed.

Secrest [205] reports the 5-year average maintenance cost for twenty-eight 25-ampere windmills on 50-foot towers was \$6.08 per year per mill. His distances between mills ranged from 1½ miles for bare pipe to 21 miles on asbestos-wrapped grease-coated pipe. A 4-month continuous record showed practically no protection 35 percent of the time. Coupons attached to the pipe show no corrosion.

# (e) STATUS OF CATHODIC PROTECTION ENGINEERING

In the light of the foregoing, it is easy to see that at present the practice of applying cathodic protection is based largely upon judgment rather than upon definitely established engineering principles. The application of principles established by laboratory experiment is very difficult because the conditions encountered in practice are complex and difficult to determine. For example, the laboratorian has definite control of the composition, concentration, and homogeneity of the solutions he uses, and can easily arrange his experiments so that the electrodes employed have equipotential surfaces. Also, the supply of oxygen and the circulation of the electrolyte are under control. No such propitious state of affairs exists in nature. The corrosion engineer has to contend with variations in composition, concentration, and aeration, and in addition with limited accessibility for the measurement of currents and potentials. He also has to take into consideration certain time effects. When an electromotive force is applied between a pipe and an anode, the potential of the pipe changes gradually for hours or days and when the current is cut off the pipe is not immediately restored to its original potential. Furthermore, the relatively large currents and low resistances encountered in field work and the impossibility of isolating the soil and pipe with which the test is concerned from the adjacent soil and pipe make precise measurements in the field both difficult and time consuming.

The immediate problem of the cathodic protection engineer has been to effect a substantial reduction in losses due to corrosion in the shortest possible time, and there has been little time or equipment available for fundamental research. It is not surprising, therefore, that the criteria for judging whether or not complete protection has been achieved are not yet well established or that many reports are somewhat indefinite as to methods and results. It is generally agreed, however, that cathodic protection is an effective method of reducing corrosion and worthy of continued development. There is still much to be learned of the underlying phenomena involved in the corrosion of buried metal and of the practical conditions to be met in the field. From the nature of the problem, it is evident that the combined efforts of research workers and engineers will be required to bring about a satisfactory solution. No doubt, the problem will be attacked more intensively than is now possible as soon as the war is over.

#### XV. REFERENCES

- [1] Second report of the Dutch Corrosion Committee II for the study of the corrosive effects of soils on pipes: Action of soil on pipes (1935).
- [2] Report by the Dutch Corrosion Committee II for the study of the corrosive effects of soils on pipes: Coating of pipe with bituminous compositions (undated).
- [3] Third report of the Dutch Corrosion Committee II for the study of the corrosive effect of soils on pipes: Regulations for the asphalting of pipes with asphaltic bitumen of the blown type (1937).
- [4] C. A. H. von Wolzogen Kuhr and L. S. van der Vlugt, The gratifization of cast iron as an electrobiological process in anaerobic soils, Water (Dutch) 18, 147 (1934).
- [5] H. Van der Veen, The investigation of the covering property of asphaltic coatings, Fourth NBS Underground Corrosion Conference (1937).
- [6] C. M. Wichers, Corrosie van gegoten ijzeren buizen in den grond, Groningen, Rapport I (1932); II (1932); III (1935); IV (1937).
- [7] C. M. Wichers, The corrosion of pipe lines due to earth contact, Fourth NBS Underground Corrosion Conference (1937). 814786°

- [8] G. O. Thomas, Fundamental requirements of pipe coatings, Commonwealth Eng. 24, 293 (Australia, 1937).
- [9] R. W. Parkhurst, Bituminous protective coatings for steel pipe, Commonwealth Eng. 24, 125 and 156 (Australia, 1936).
- [10] G. O. Thomas. Determination of the suitability of bituminous coatings for underground use, Inst. Engrs. 6, 337 (Australia, 1934).
- [11] C. C. Challis, Mitigation of corrosion on the distribution system of the Australian Gas Light Co., Fourth NBS Underground Corrosion Conference (1937).
- [12] G. O. Thomas, The measurement of water absorption as a criterion of the protective properties of protective coating materials, Fourth NBS Underground Corrosion Conference (1937).
- [13] C. M. Longfield, Stray current electrolysis in Australia, Fourth NBS Underground Corrosion Conference (1937).
- [14] R. J. Dumas, Description of outer pipe coating, Fifth NBS Underground Corrosion Conference (1943).
- [15] G. O. Thomas, Manufacture of coal tar enamel pipe coatings, Fifth NBS Underground Corrosion Conference (1943).
- [16] J. C. Hudson, T. A. Banfield, and H. A. Holden, Tests on the corrosion of buried ferrous metals, Iron & Steel Inst. Paper No. 6 (England, 1942).
- [17] H. J. Bunker, Micro-biological experiments in anaerobic corrosion, J. Soc. Chem. Ind. 58, 93 (England, 1939).
- [18] W. G. Radley, Determination of the causes of sheath corrosion, Elec. Engrs. 57, 168 (1938).
- [19] Report of the American Committee on Electrolysis, Am. Inst. Elec. Engrs. (New York, N. Y., 1921).
- [20] U. R. Evans, Metallic corrosion, passivity and protection (Edward Arnold & Co., London, 1937).
- [21] O. Krohnke, E. Maas, and W. Beck, Die Korrosion, Band 1 (Verlag von S. Hirzel, Leipzig, 1929).
- [22] J. W. Shipley, The corrosion of cast iron and lead pipes in alkaline soil, J. Soc. Chem. Ind. 41, 311 (1922).
- [23] J. W. Shipley and I. R. McHaffie, The graphite softening of cast iron, Ind. & Chem. Eng. 16, 573 (1924).
- [24] J. W. Shipley, I. R. McHaffie, and N. D. Clare, Corrosion of iron in the absence of oxygen, Ind. & Chem. Eng. 17, 381 (1925).
- [25] J. W. Shipley and I. R. McHaffie, The relation of hydrogen ion concentration to the corrosion of iron, Chem. & Met. 8, 121 (Canada, 1924).
- [26] W. Nelson Smith, The principles of three-wire distribution for electric railways, Eng. J. 6 (No. 5) 235 (1923).
- [27] E. P. Fetherstonhaugh, Studies of cast iron pipe corrosion, Fourth NBS Underground Corrosion Conference (1937).
- [28] F. N. Speller, Corrosion, causes and prevention (McGraw-Hill Book Co., New York, N. Y., 1935).
- [29] S. P. Ewing, Soil corrosion and pipe line protection, Am. Gas Assn. (New York, N. Y., 1938).
- [30] R. M. Burns and A. E. Schuh, Protective coatings for metals (Reinhold Publishing Corporation, New York, N. Y., 1938).
- [31] T. P. Hoar, The principles of metallic corrosion, Science of petroleum, p. 2307 (Oxford University Press, England, 1938).
- [32] O. Gatty and E. C. R. Spooner, The electrode potential behavior of corroding metals in aqueous solutions (The Clarendon Press, Oxford, 1938).
- [33] C. F. Marbut, Atlas of American agriculture, part III, Soils of the United States (U. S. Government Printing Office, 1935).
- [34] Mark Baldwin, Chas. E. Kellogg, and James Thorp, Soil classification; Soils & Men, p. 979 (U. S. Government Printing Office, 1938).
- [35] Kirk H. Logan, Engineering significance of National Bureau of Standards soil corrosion data, J. Research NBS 22, 109 (1939) RP1171.

- [36] J. R. Baylis, Prevention of corrosion and red water, J. Am. Water Works Assn. 15, 596 (1926).
- [37] W. G. Whitman, R. P. Russell, and V. J. Altieri, Effect of hydrogen-ion concentration on submerged corrosion of steel, Ind. Eng. Chem. 16, 665 (1924).
- [38] W. G. Whitman and R. P. Russell, The submerged corrosion of iron, Soc. Chem. Ind. 43, 193T (1924).
- [39] I. A. Denison and R. B. Hobbs, Corrosion of ferrous metals in acid soils, J. Research NBS 13, 125 (1934) RP696.
- [40] I. A. Denison and S. P. Ewing, Corrosiveness of certain Ohio soils, Soil Sci. 40, 287 (1935).
- [41] G. N. Scott, The use and behavior of protective coatings on underground pipes, Proc. Am. Petroleum Inst. [I] 10, 78 (1929).
- [42] B. A. Keen, The Physical Properties of the Soils (Longmans Green & Co., New York, N. Y., 1931).
- [43] T. D. Rice and L. T. Alexander, The physical nature of the soil, Soils & Men, p. 887 (U. S. Government Printing Office, 1938).
- [44] H. G. Byers, M. S. Anderson, and Richard Bradfield, General chemistry of soils, Soils & Men, p. 911 (U. S. Government Printing Office, 1938).
- [45] K. H. Logan, Soil-corrosion studies, 1934. Rates of loss of weight and pitting of ferrous specimens, J. Rosearch NBS 16, 431 (1936) RP883.
- [46] Soil Survey of Baltimore County, Md. (U. S. Government Printing Office, 1919).
- [47] Melvin Romanoff, Effect of aeration on the hydrogen-ion concentration of soils in relation to the identification of corrosive soils, J. Research NBS 34, 227 (1945) RP1630.
- [48] K. H. Logan, Soil-corrosion studies, 1937. Corrosion-resistant materials and special tests, J. Research NBS 23, 515 (1939) RP1250.
- [49] E. R. Shepard, Some factors involved in soil corrosion, Ind. Eng. Chem. 26, 723 (1934).
- [50] G. N. Scott, Adjustment of soil corrosion pit depth measurements for size of sample, Proc. Am. Petroleum Inst. [IV] 14, 204 (1934).
- [51] E. P. Fetherstonhaugh, Discussion of underground corrosion, Proc. Am. Soc. Civil Engrs. 101, 828 (1936).
- [52] J. F. Putnam, Soil corrosion, Proc. Am. Petroleum Inst. [IV] 16, 66 (1935).
- [53] J. F. Brennan, A mathematical theory of corrosion, Gas Age-Record 75, 359 (1935).
- [54] G. Corfield, Running a soil survey on a large distribution system, Western Gas, 6 (No. 3) 25 (1930).
- [55] G. N. Scott, A preliminary study of the rate of pitting of iron pipe in soils, Proc. Am. Petroleum Inst. [IV] 14, 212 (1934).
- [56] K. H. Logan, S. P. Ewing, and I. A. Denison, Soil corrosion testing, Symposium on Corrosion Testing Procedures, Am. Soc. Testing Materials, p. 95 (1937).
- [57] K. H. Logan and M. Romanoff, Soil-corrosion studies, 1941: Ferrous and nonferrous corrosion-resistant materials and nonbituminous coatings, J. Research NBS 33, 145 (1944) RP1602.
- [58] Report on steel pipe lines for underground water service, Underwriters Laboratories, Special Investigation 888, Appendix IV (1936).
- [59] Standard for steel pipe lines for underground water service, Underwriters Laboratories, Special Investigation 888—38 (1938).
- [60] K. H. Logan, W. Rogers, and J. F. Putnam, Pipe line currents, Am. Petroleum Inst. Production Bul. 204, 116 (1930).
- [61] K. H. Logan, API pipe-coating tests—Final report, Proc. Am. Petroleum Inst. [IV] 21, 32 (1940).
- [62] K. H. Logan, Soil corrosion studies, 1934. Rates of loss of weight and penetration of nonferrous materials, J. Research NBS 17, 781 (1936) RP945.
- [63] R. J. McKay and R. Worthington, Corrosion resistance of metals & alloys (Reinhold Publishing Corporation, New York, N. Y., 1936).

- [34] K. H. Logan, Soil corrosion studies 1939. Ferrous and nonferrous corrosionresistant materials, J. Research NBS 28, 379 (1942) RP1446.
- [65] W. W. H. Gee, Electrolytic methods for preventing corrosion, Trans. Faraday Soc. 9, 115 (1916).
- [66] K. H. Logan and S. P. Ewing, Soil corrosion studies 1934. Field tests of nonbituminous coatings for underground use, J. Research NBS 18, 361 (1937) RP982.
- [67] H. S. Rawdon, Protective metallic coatings (The Chemical Catalog Co., New York, N. Y., 1938).
- [68] R. V. A. Mills, Protection of oil and gas fuel equipment against corrosion, U. S. Bureau of Mines Bul., 233 (1925).
- [69] K. H. Logan, S. P. Ewing, and C. D. Yeomans, Bureau of Standards soil-corrosion studies I. Soils, materials and results of early observations, Tech. Pap. BS 22, 447 (1928) T368.
- [70] C. O. Bannister, The protective action of zinc plates in boilers, Metal Ind. 41, 441-43, 467-70 (London, 1932).
- [71] P. D. Merica, Structure of the coating on tinned sheet copper in a specific case of corrosion, Tech. Pap. BS 22 (1917) T90.
- [72] B. M. McCollum and O. S. Peters, Surface insulation of pipes as a means of preventing electrolysis, Tech. Pap. BS 2 (1914) T15.
- [73] G. N. Scott, API pipe coating tests: Progress reports I and II, Proc. Am. Petroleum Inst. [IV] 12, 55, 72 (1931); Progress report III, Proc. Am. Petroleum Inst. [IV] 13, 114 (1932); Progress report IV, Proc. Am. Petroleum Inst. [IV] 15, 18 (1934).
- [74] E. R. Shepard, Measurement of the electrical conductance of nonmetallic coatings, Am. Gas J. 136, 22 (1932).
- [75] S. P. Ewing, Field tests of protective coatings, Am. Gas Assn., Proc. p. 627 (1936).
- [76] J. F. Putnam, Private communication.
- [77] K. H. Logan, The effect of protective coatings on the rate of pitting of pipe lines, Proc. Am. Petroleum Inst. [IV] 22, 34 (1941).
- [78] R. F. Hadley, Studies in microbiological anaerobic corrosion, Am. Gas Assn., Proc. p. 1764 (1940).
- [79] T. D. Beckwith, The bacterial corrosion of iron and steel, Am. Water Works Assn. 33, 147 (1941).
- [80] I. A. Denison, Electrolytic measurements of the corrosiveness of soils, J. Research NBS 17, 363 (1936) RP918.
- [81] I. A. Denison, Making soil-corrosion survey by using electrolytic test, Oil & Gas J. 37 (No. 19) 96 (1938).
- [82] A. Hickling, Studies in electrode polarization. Part I—The accurate measurement of the potential of a polarized electrode, Trans. Faraday Soc. 33, 1540 (1937).
- [83] R. Darnielle, Measurement of electrode potentials and polarization in soil corrosion cells, J. Research NBS 25, 421 (1940) RP1336.
- [84] L. C. Bannister and U. R. Evans, Passivity of metals, part V, the potentialtime curves of some iron alloys, J. Chem. Soc. p. 1361 (1930).
- [85] R. M. Burns, Corrosion of metals—I. Mechanism of corrosion processes, Bell System Tech. J. 15, 20 (1936).
- [86] R. H. Brown and R. B. Mears, Application of electrochemical measurements to studies of the corrosion of 18-8 stainless steel, Trans. Faraday Soc. 35, 467 (1939).
- [87] U. R. Evans, The distribution and velocity of the corrosion of metals, J. Franklin Inst. 208, 45 (1929).
- [88] U. R. Evans and T. P. Hoar, The velocity of corrosion from the electrochemical standpoint, part II, Proc. Roy. Soc. [A] 137, 343 (1932).
- [89] W. J. Müller, The effect of cathodic reaction on the corrosion of metals from the viewpoint of the local cell theory, Trans. Electrochem. Soc. 79, 169 (1939).

- [90] O. Gatty and E. C. R. Spooner, The electrode potential behavior of corroding metals in aqueous solutions (Clarendon Press, Oxford, England, 1938).
- [91] R. B. Mears, Discussion, Trans. Electrochem. Soc. 69, 164 (1936).
- [92] U. R. Evans, L. C. Bannister, and S. C. Britton, Velocity of corrosion from the electrochemical standpoint, Proc. Roy. Soc. [A] 131, 367 (1931).
- [93] Scott Ewing, Determination of the current required for cathodic protection, Am. Gas Assn. Proc. op. 613 (1940).
- [94] I. A. Denison and R. B. Darnielle, Observations on the behavior of steel corroding under cathodic control in soils, Trans. Electrochem. Soc. 76, 199 (1939).
- [95] I. A. Denison, Electrolytic behavior of ferrous and nonferrous metals in soil corrosion circuits, Trans. Electrochem. Soc. 81, 435 (1942).
- [96] T. P. Hoar, The electrochemistry of protective metallic coatings, J. Electrodepositors' Tech. Soc. 14, 33 (1938).
- [97] E. R. Shepard, Pipe line currents and soil resistivity as indications of local corrosive soil areas, BS J. Research 6, 683 (1931) RP298.
- [98] F. Wenner, A method of measuring with resistivity, BS Sci. Pap. 12, 469 (1916) S258.
- [99] W. J. Rooney, Earth resistivity measurements in the copper country, Michigan, Terrestrial Magnetism and Atmospheric Electricity 32, 97 (1927).
- [100] Ground resistance testing, Tech. Bul. 1285 (James G. Biddle Co., Philadelphia, Pa., 1931).
- [101] H. W. Wahlquist, Private communication.
- [102] R. O. Davis, The use of the electrolytic bridge for determining soluble salts, U. S. Dept. Agr. Circular 423 (1927).
- [103] Wm. E. Huddleston, The value of the radio balance in conducting soil surveys, Petroleum Ind. Elec. News 11 (No. 10) 53 (1942).
- [104] B. B. Legg, Early steps in the development of a corrosivity apparatus, Natural Gas 12 (No. 2) 10 (1931).
- [105] R. H. Hadley, Pipe coatings and corrosion research, Gas Age 92, 29 (Dec. 2, 1943).
- [106] Burton McCollum and K. H. Logan, Earth resistivity and its relation to electrolysis of underground structures, National Bureau of Standards Tech. Pap. 23 (1916).
- [107] H. D. Holler, Corrosiveness of soils with respect to iron and steel, Ind. Eng. Chem. 21, 750 (1929).
- [108] I. A. Denison, Methods for determining the total acidity of soils, J. Research 10, 413 (1933) RP539.
- [109] S. P. Ewing, Soil corrosion and pipe line protection (Am. Gas. Assn., 1938).
- [110] J. F. Putnam, Electrolysis, Sibley J. Eng. 31, 88 (1917).
- [111] K. H. Logan and E. A. Koenig, A comparison of methods for estimating the corrosivity of soils, Oil and Gas J. 38 (No. 27) 130 (1939).
- [112] C. R. Weidner and L. E. Davis, Relation of pipe line currents and soil resistivity to corrosion, Proc. Am. Petroleum Inst. [IV] 12, 36 (1931).
- [113] C. Fitzgerald, Discussion of relation of pipe line currents and soil resistivity to corrosion, by C. R. Weidner and L. E. Davis, Am. Petroleum Inst. [IV] 12, 47 (1931).
- [114] Stanley Gill and Walter Rogers, Relation of long-line currents to corrosion, Physics 1, 194 (1931).
- [115] Scott Ewing, Rough correlation between corrosiveness and resistivity for alkali soils, Oil and Gas J. 30, 29 (1932).
- [116] K. H. Logan, W. F. Rogers, and J. F. Putnam, Pipe line currents, Oil and Gas J. 28 (No. 29) 130 (1930).
- [117] J. C. Sterling, Pipe line currents, Oil and Gas J. 37 (No. 19) 142 (1938).
- [118] C. H. McRaven, Measurements of pipe line currents, Petroleum Ind. Elec. News (April 1941).
- [119] Carl Hering, Measurement of stray electric currents in underground pipes, Trans. Am. Inst. Elec. Engrs. 31, 1449 (1912).

- [120] O. C. Mudd, Detecting pipe line corrosion with electrical devices, Oil and Gas J. 38 (No. 5) 48 (1939).
- [121] O. C. Mudd, Use of soil surface potentials in locating pipe line corrosion, Oil and Gas J. 41 (No. 1) 50 (1942).
- [122] C. M. Schlumberger and E. G. Leonardon, Location and study of pipe line corrosion by surface electrical measurements, Am. Inst. Mining Met. Engrs. Tech. Pub. No. 476 (1932).
- [123] J. M. Pearson, Electrical instruments applied to the study of pipe line corrosion, Proc. Am. Petroleum Inst. [IV] 16, 75 (1935).
- [124] B. McCollum and K. H. Logan, Electrolysis testing, Tech. Pap. BS (1927) T355.
- [125] B. McCollum and K. H. Logan, Practical applications of the earth current meter, Tech. Pap. BS 21, 683 (1927) T351.
- [126] W. R. Schneider, Corrosion coupons and pipe life predictions, Fifth NBS Underground Corrosion Conference (Abstract), Gas 20 (No. 4) 39 (1944).
- [127] Valuation Docket 1203, Interstate Commerce Commission (1937).
- [128] Stanley Gill, Locating pipe line inspection points, Oil Weekly (May 30, 1923).
- [129] K. H. Logan and A. E. Koenig, Methods of inspecting pipe lines, J. Am. Water Works Assn. 31, 1451 (1939).
- [130] D. F. Van deWater, Preventive maintenance by systematic pipe line inspection, Petroleum Eng. 14 (No. 8) 74 (1943).
- [131] G. N. Scott, The use and behavior of protective coatings on underground pipes, Am. Petroleum Institute Bul. 10 (No. 2) 78 (1929); Oil and Gas J. 27 (No. 29) 127 (1929).
- [132] S. P. Ewing and G. N. Scott, An electrolytic method for detecting the condition of a non-metallic pipe coating, Tenth annual Am. Gas Assn. Conference (1933).
- [133] G. W. Clarvoe, The detection of flaws in pipe line protective coatings before burial, Pipe Line News 5 (No. 8) 13 (1933).
- [134] D. Harrell, Detection of holidays in pitch enamel coatings, Petroleum Eng. 9, 97 (1936).
- [135] D. E. Stearns, Accurate voltage control was a feature of new holiday detector, Petroleum Eng. 12, 160 (1941).
- [136] D. E. Stearns, Electronic holiday detector provides robot pipe inspection service, Am. Gas Assn. Monthly 26 (No. 3) 115 (1944).
- [137] J. M. Pearson, Electrical examination of coatings on buried pipe, Petroleum Eng. 12, 82 (1941).
- [138] G. N. Scott and S. P. Ewing, Pipe line fabrics, Am. Dyestuff Reporter 24, 699 (1935).
- [139] S. P. Ewing, Studies of coatings for pipe lines, Am. Gas Assn. Proc., p. 774 (1931).
- [140] V. A. Grodsky, A laboratory soil corrosion test, Am. Water Works Assn. 30, 760 (1938).
- [141] C. M. Wichers, The corrosion of pipe lines due to the earth contact, Fourth NBS Underground Corrosion Conference (1937). Unpublished.
- [142] T. F. P. Kelly, Performance of coal tar protective coatings, Gas 20 (No. 6) 31 (1944).
- [143] J. H. Peper, Use of cement in protecting underground pipe, Oil and Gas J. 32, (No. 33) 9; (No. 37) 34 (1934).
- [144] L. E. Bredberg, Reconditioning lines in salt and marsh districts, Oil and Gas J. 33 (No. 3) 8 (1933).
- [145] Standard specifications for coal 'tar enamel protective coatings for steel water pipe, Am. Water Works Assn. specification 7A. 5 (1940) 7A. 6 (1940).
- [146] F. H. Love, Constructing the large diameter war emergency pipe line, Petroleum Eng. 14, 39 (1943).
- [147] F. H. Love, Constructing the War Emergency Pipe products line, Petroleum Eng. 15, 133 (1944).

- [148] J. C. Sterling, Field application of pipe line coatings, Fourth NBS Underground Corrosion Conference (1937). Unpublished.
- [149] G. O. Thomas, Manufacture of coal tar enamel pipe coatings, Fifth NBS Underground Corrosion Conference (1943). Unpublished.
- [150] W. F. Rogers, Performance data on enamel type pipe line coatings. Petroleum Eng. 14, 162 (1943).
- [151] R. F. Hadley, Private communication.
- [152] L. A. Hugo, Experience in the use of the Pearson Electronic Coating Inspector (Abstract), Oil and Gas J. 42 (No. 51) 50 (1944).
- [153] G. B. McCabe and F. M. Hull, Detroit pioneers with pipe type gas filled cable, Elec. World 117 (No. 26) 2214 (1942).
- [154] L. F. Sherer, Cooperative problems in cathodic protection, Oil and Gas J. 38 (No. 37) 179 (1939).
- [155] G. I. Rhodes, Cathodic protection or electrical drainage of bare pipe lines (Monograph), Am. Gas Assn. (1935).
- [156] W. Ryland Hill, Laboratory tests of cathodic protection of steel in various corrosive solutions, Petroleum Eng. 12 (No. 13) 51 (1941).
- [157] W. F. Rogers, Relationship of current density to cathodic protection, Petroleum Eng. 12 (No. 1) 156 (1940).
- [158] J. H. Keeling, Current and voltage needs for protecting steel submarine pipe lines, Gas 15 (No. 9) 31 (1939).
- [159] G. I. Rhodes, Two unusual installations of cathodic protection, Proc. Am. Petroleum Inst. 17 (No. 4) 21 (1936).
- [160] O. C. Mudd, Locating pipe line corrosion by soil surface potential measurements, Petroleum Ind. Elec. News 11 (No. 11) 17 (1942).
- [161] R. J. Kuhn, Cathodic protection of underground pipe lines from underground corrosion, Proc. Am. Petroleum Inst. [IV] 14, 153 (1933).
- [162] D. Holsteyn, Practical design and economics of a cathodic unit as applied in the refinery, Petroleum Ind. Elec. News 13 (No. 3) 9 (1943).
- [163] R. A. Brannon, Cathodic protection of tank farms, Petroleum Ind. Elec. News 12 (No. 6) 11 (1942).
- [164] S. P. Ewing, Cathodic protection of pipe lines from soil corrosion, Gas Age-Record 75 (No. 9) 179, (No. 10) 219, (No. 11) 239, (No. 12) 261 (1935); Natural Gas 16 (No. 3) 5, (No. 4) 16 (1935).
- [165] R. J. Kuhn, Galvanic currents on cast iron pipe, First NBS Underground Corrosion Conference (1928). Unpublished. (Abstract) Ind. Eng. Chem. 22, 335 (1930).
- [166] G. N. Scott, An analysis of certain circuits in cathodic protection, Proc. Am. Petroleum Inst. [IV] 23, 36 (1942).
- [167] G. N. Scott, A rational approach to cathodic protection problems, Petroleum Eng. 12 (No. 8) 271 (1941).
- [168] F. P. Bowden, The effects of pH on overpotential, Trans. Faraday Soc. 24, 473 (1928).
- [169] J. M. Pearson, Null methods applied to corrosion measurements, Trans. Electrochem. Soc. 81, 485 (1942).
- [170] K. H. Logan, Determination of current required for cathodic protection, Petroleum Eng. 14 (No. 10) 168 (1943).
- [171] J. M. Pearson, Measurements of cathodic polarization and problems of interference on underground structures, Fifth NBS Underground Corrosion Conference (1943). Unpublished.
- [172] G. R. Olson, Control of pipe line corrosion, Petroleum Eng. 14 (No. 12) 96 (1943).
- [173] U. S. Patent 1962696 (1934).
- [174] S. U. McGary, Determining the location and capacity of units for cathodic protection, Petroleum Eng. 9 (No. 11) 40 (1938).
- [175] W. F. Rogers, Methods of designing cathodic protection installations, Petroleum Eng. 12 (No. 6) 100 (1941).
- [176] Starr Thayer, The development and application of electrical protection for pipe lines, Proc. Am. Petroleum Inst. [IV] 14, 143 (1933).
  [177] W. R. Schneider, Cathodic protection of pipe lines, Gas Age-Record 71 (No. 14) 355 (1933).

- [178] G. R. Olson, Recent developments in cathodic protection of bare pipe lines, Petroleum Ind. Elect. News 12 (No. 7) 11 (1942).
- [179] R. T. Fryer, Rectifiers, all types; comparison and operation, Petroleum Ind. Elect. News 14 (No. 1) 17 (1944).
- [180] A. C. Aliter, The cathodic protection of bare steel pipe in western soil types by the use of zinc or aluminum, Convention Pacific Coast Gas Association (1941).
- [181] G. R. Olson, A field experiment with magnesium rods, Proc. Nat. Assn. Corrosion Engrs. Convention 1, 56 (1944).
- [182] J. J. Grebe and R. E. McNulty, Magnesium and anodes in cathodic protection, Fifth NBS Underground Corrosion Conference (1943).
- [183] W. T. Smith and T. C. Marshall, Zinc for cathodic protection of pipe, Gas-Age 84 (No. 4) 15 (1939).
- [184] C. L. Brockschmidt, A practical application of zinc anode protection to an 18 inch pipe line, Petroleum Ind. Elec. News 11 (No. 10) 31 (1941).
- [185] O. C. Mudd, Experiences with Zinc Anodes, Petroleum Ind. Elec. News 13 (No. 1) 11 (1943).
- [186] H. W. Wahlquist, Use of zinc for cathodic protection, Proc. Nat. Assn. of Corrosion Engrs. Convention 1, 61 (1944).
- [187] W. R. Schneider, Pilot grounds for cathodic protection of pipe lines, Western Gas 10 (No. 8) 14 (1934).
- [188] O. S. Peters, Ground connections for electrical systems, Tech. Pap. BS (1918) T108.
- [189] O. H. Gish and W. J. Rooney, Measurement of resistivity of large masses of undisturbed earth, Terrestrial Magnetism and Atmospheric Electricity 30, 161 (1925).
- [190] B. McCollum and K. H. Logan, Electrolytic corrosion of iron in soils, Tech. Pap. BS (1914) T25.
- [191] O. C. Roddey and E. R. Shepard, Carbon anodes in an electrical drainage system, Fifth NBS Underground Corrosion Conference. Unpublished, (1943).
- [192) M. J. Dorcas, Ground anodes, Fifth NBS Underground Corrosion Conference (1943).
- [193] A. W. McAnney, Insoluble anodes, Petroleum Ind. Elec. News 10 (No. 3) (1940).
- [194] A. V. Smith, Cathodic interference from cathodic protection installations, Am. Gas. Assn. Monthly 25, 421 (1943).
- [195] R. J. Kuhn, Cathodic protection of pipe lines in city and country, Oil and Gas J. 36 (No. 18) 201 (1937).
- [196] W. R. Schneider, Electrical protection of city networks, Fourth NBS Underground Corrosion Conference (1937).
- [197] R. M. Wainright, Cathodic protection on distribution systems, Petroleum Ind. Elec. News 12 (No. 1) 33 1942.
- [198] Intercompany procedure report, Petroleum Ind. Elec. News 12 (No. 2) 51 (1942).
- [199] G. I. Rhodes, Electrical pipe line drainage with cost data, Elec. J. 33, 91 (1936).
- [200] D. H. Bond, Cathodic protection of oil storage tank bottoms, Petroleum Eng. 11 (No. 6) 100 (1940).
- [201] O. C. Roddey and E. R. Shepard, Distributed anode method cuts cost of cathodic protection, Oil and Gas J. 38 (No. 19) 84.
- [202] W. R. Schneider, Comparing equipment costs in cathodic protection, a digest, Gas 15 (No. 6) 31 (June 1939).
- [203] Starr Thayer, The application and economics of electrical protection of pipe lines, Proc. Am. Petroleum Inst. [IV] 17 (No. 12) 33 (1936).
- [204] W. H. Stewart, Problems in connection with protection of bare pipe, Petroleum Ind. Elec. News 13 (No. 2) 17 (1943).
- [205] L. C. Secrest, Cathodic protection—its application to a pipe line, Oil and Gas J. 43 (No. 3) 82 (1945).
- [206] R. K. Schofield, The pF of water in soil, Trans. Third Intern. Cong. Soil Sci. 2, 37 (England, 1935).

# XVI. APPENDIX 1. COOPERATORS WITH THE NATIONAL BUREAU OF STANDARDS IN THE CORROSION INVESTIGATIONS

# 1. FURNISHERS OF LABOR AND OF TEST SITES

Works. Atlanta Gas Light Co. Atlantic City Gas Co. Atlantic Pipe Line Co. City of Baltimore, Department of Public People's Water & Gas Co. Boston Consolidated Gas Co. Brockton Gas Light Co. Camden Water Department. Carolina Power & Light Co. Cincinnati Gas & Electric Co. City of Charleston, Water Department. The Citadel. City of Cleveland, Department of Public Ŭtilities. Colorado Interstate Gas Co. Community Natural Gas Co. Dallas Gas Co. Delaware Power & Light Co. Des Moines Gas Co. Duke Power Co. East Bay Municipal Utility District. East Ohio Gas Co. Empire Pipe Line Co. Equitable Gas Co. Florida Power & Light Co. Florida Public Utilities Co. Georgia Railway & Power Co. Gulf Oil Corporation. Humble Pipe Line Co. Indiana Pipe Line Co. City Commission of Jacksonville. Jacksonville Gas Co. ties. Kansas City Gas Co. Los Angeles Gas & Electric Co. Macon Gas Co. Memphis Board of Water Commissioners. City of Meridian. City of Middleboro, Mass. Midwest Refining Co. City of Milwaukee, Dept. of Public Works. Milwaukee Gas Light Co. Mississippi Power Co. Mississippi River Fuel Corporation. City of Mobile, Water Works Department. Mountain Fuel Supply Co.

New Orleans Audubon Park Commission. New Orleans Sewerage & Water Board.

New Orleans Public Service, Incorporated.

North Carolina Public Service Co.

Northern States Power Co.

Albuquerque Gas & Electric Co.

Alexandria Water Co.

City of Norwood, Mass. Ohio Fuel Gas Co. City of Atlanta, Department of Water Oklahoma Pipe Line Co. The Omaha and Council Bluffs Electrolysis Committee. Pacific Gas & Electric Co. Pensacola-Gulf Power Co. Philadelphia Electric Co. City of Phoenix, Water Department. Prairie Pipe Line Co. Public Service Company of Colorado. Pueblo Gas & Fuel Co. Pure Oil Pipe Line Co. Raleigh Gas Co. City of Rochester, Dept. of Public Works. Rochester Gas & Electric Company. San Antonio Public Service Board. San Diego Consolidated Gas & Electric San Joaquin Light & Power Corporation. City of Seattle, Department of Public Works. Shell Oil Co. Shell Petroleum Corporation. Shell Pipe Line Corporation. Sinclair Pipe Line Co. Sinclair-Prairie Pipe Line Company of Texas. Sohio Pipe Line Co. Southern California Gas Co. Southern California Telephone Co. Southern Cities District Co. Southern Natural Gas Co. Southwestern Gas & Electric Co. City of Kalamazoo, Dept. of Public Utili- City of Springfield, Ohio, Water Department. Springfield Railway Co. Standard Oil Company of California. Standard Oil Company of Louisiana, Pipe Line Dept. Standard Oil Development Co. Standard Oil Company of New Jersey. Stanolind Oil & Gas Co. Sun Oil Line Co. Susquehanna Pipe Line Co. Tampa Gas Co. Tidal Pipe Line Co. Tidewater Pipe Line Co., Ltd. Tri City Railway & Light Co. United Gas Pipe Line Co. The United Light & Power Service. Union, Light, Heat & Power Co. Union Light & Railway Co. Union Oil Company of California. Vicksburg Gas Co. Wilmington Gas Co.

## 2. SUPPLIERS OF MATERIALS

#### (a) FERROUS

Allegheny Ludlum Steel Corporation.
American Cast Iron Pipe Co.
American Radiator Co.
American Rolling Mill Co.
H. A. Brassert Co.
A. M. Byers Co.
Carnegie-Illinois Steel Corporation.
Carson Cadillae Corporation.
The Duriron Co., Inc.
Electric Steel Founders.
Electro Metallurgical Co.
Inland Steel Co.
The International Nickel Co., Inc.
Jones and Laughlin Steel Corporation.
Lukenheimer Co.

McWane Cast Iron Pipe Co.

Mechanite Metals Co.
National Cast Iron Pipe Co.
National Tube Co.
National Tube Co.
Nugent Steel Casting Co.
Pittsburgh Valve, Foundry & Construction Co.
Reading Iron Co.
Republic Steel Corporation.
Sharon Steel Corporation.
Sivier Steel Castings Co.
Stockham Pipe & Fittings Co.
Union Carbide & Carbon Research Laboratories.
United States Pipe & Foundry Co.
Walworth Co.

(b) NONFERROUS

American Brass Co.
American Smelting & Refining Co.
American Zinc Products Co.
Bell Telephone Laboratories Inc.
Bridgeport Brass Co.
Brown Co.
Chase Brass & Copper Co.
General Cable Corporation.
Habirshaw Electric Cable Co.
The Hoyt Metal Co.
Y. T. Hungerford Brass & Copper Co.
Illinois Zinc Co.

Johns-Manville Sales Corporation.
Keasby & Mattison Co.
Lumen Bearing Co.
Mueller Brass Co.
Mueller Co.
National Carbon Co.
National Lead Co.
The New Jersey Zinc Co.
Revere Copper & Brass, Incorporated.
Scovill Manufacturing Co
Sharon Steel Hoop Co.
Standard Underground Cable Co.

Youngstown Sheet & Tube Co.

## (c) COATINGS

Albrecht Pagenstecher. Aluminum Company of America. American Machine & Foundry Co. American Tar Products Co. Arco Co. The Bakelite Co. Ball Chemical Co. Barber Asphalt Co. The Barrett Co. Benjamin Foster Co. Calorizing Co. Chadeloid Chemical Co. Consolidated Gas Company of New York. Dearborn Chemical Co. E. I. du Pont de Nemours & Co., Inc. Eagle-Picher Lead Co. Emulsion Process Co. Ferro Enamel Corporation. Fish-Schurman Corporation. The Flintkote Co. General Paint Corporation. The P. D. George Co. The B. F. Goodrich Rubber Co. Goodyear Tire & Rubber Co. Harpoon Paint Products, Incorporated.

Headley Emulsified Products Co. Hill, Hubbell, & Company. Inertol Co. Iroquois Gas Corporation. Irvington Varnish & Insulator Co. The Locomotive Terminal Improvement Co. McEverlast, Inc. Andrew McLean Co. Merchants Basket & Box Co. Paraffine Companies, Inc. Pioneer Asphalt Co. Resisteor Engineering Corporation. H. H. Robertson Co. Shori Process Corporation. Sherwin-Williams Co. Jas. B. Sipe & Co. Southport Paint Co., Inc. Technical Products, Inc. The Texas Co. Thickol Corporation. Udylite Process Co. United Gas Impressment Co. Wailes Dove-Hermiston Corporation.

## 3. ADVISORY ORGANIZATIONS

American Committee on Electrolysis, Re- | American Standards Association. search Subcommittee. American Engineering Standards Committee. American Foundrymen's Association. American Gas Association. American Petroleum Institute. American Society for Testing Materials.

American Water Works Association. American Zinc Institute, Inc. Asphalt Institute. Cast Iron Pipe Research Association. Copper and Brass Research Association. Lead Industries Association.

#### 4. GOVERNMENT DEPARTMENTS

Department of Agriculture, Bureau of Plant Industry.

Department of Commerce, National Bupartment o reau of Standards.

# XVII. APPENDIX 2. DESCRIPTIONS OF SOILS AT TEST SITES

Prepared by M. Romanoff. The profiles lave been described by S. Ewing, I. A. Denison, G. N. Scott, and by the following soil surveyors from the Bureau of Plant Industry of the United States Department of Agriculture: A. E. Taylor, M. H. Laphan, R. Wildermuth, W. J. Geib, H. H. Bennett, H. G. Lowis, F. A. Hayes, W. T. Carter, R. C. Roberts, Mark M. Baldwin, R. S. Emith.
When the profile at the test side was not described the typical profile of the soil type was taken from soil-survey reports.

Table 101.—National Bureau of Standards test sites.

	Depth of speci- mens	Inches	96		Below 40	8
-	Topography		Undulating to gently roll- ing.		do	Moderateslope.
	Internal drainage		Poor		op	Little exes-
	Description of sail profile	0-8 grayish yellow or yellowish gray silt hoam mottled with yel- low and yellowish brown. 8-23 mottled yellow and gray silty clay loam which contains frag- ments of shale.	23-30 bluish gray silty alay loam with bands of yellow indicating the bedding planes of the shale.  370 silty calky or silty clay hom layer of shale which has a bluish gray of or and is attended along bedding planes with yellow.  70-90 reddish brown shale streaked with gray.  76-90 compact bluish gray shale with yellowish brown and reddish	orown ferans. 96-100 the streaks become less conspicuous. This shale runs high in aluminum sulfate, which, with water, breaks down into aluminum hydroxide and sulfurie acid.	(0-10 black to dark brown silty clay 10-10 black to dark brown silty clay 10-740 black clay. No definitely residual matter was discovered within 40 inches. Small rounded quartatic gravel and line concentrations disseminated through the subsoil.	0-8 grayish brown, rather compact, very fine sandy loam. A few fragments of granite and quartz found on the surface.  8-10 transition layer into.  10-28 compact brittle red day containing very few mice flakes and practically no stand and stones.  32-70 miraceous, more friable, and not as compact as above horison, red day foam or eds.  48-82 layer of sandy clay with yellowish mottlings.  25-77 or an infraceous day as in 32-84.  74-84 end very fine sandy loam with yellowish mottlings crystals, and having a few brownish and yellowish mottlings of yellowing a few brownish and yellowish mottlings due to partially decomposed rock.
	Location		:		Dallas, Tex	Atlanta, Ga
	Soil type		Allis silt loam		Bell clay	Cecil clay loam
	Site No.		-		61	က

98	30	36	55
Gently rolling	Smooth and level.	Moderately rolling.	Smooth ridge top.
poog	Poor	Excessive	Pair
0-10 grayish brown mollow loam gradually getting lighter in color with inneressing depth.  The top 6 inches of the tench is a mixture of road material and sail. No vegetlation, tench is a mixture of road material and 10-34 mellow, only slightly darker in color and heavier in exture with increasing depth.  34-46 minereous rather loose frable siltloam containing considerable meant. At 36 inches there is a layer of partially decomposed grante.  Sail in this site is considerably wetter than the average condition of this soil, as the tranch goes all the rain water that fails on the adjacent highway.	0–10 dark dull gray or drab elay of adobe structure, siteky when wet, contain numerous plants and grass roots and an appreciable amount of fine gritty material and gravel fragments.  1-63 eligithy more compact brownish gray or drab friable elay which is sticky when wet.  which is sticky when wet.  It contains spherical shotlike iron concretions of black streaks. It contains spherical shotlike iron concretions of black toor, rasquign is as from a phich ad to rabin black soot, rasquign is as from a phich ad to rabin black soot.  30–48 soil grades into a yellowish brown sily clay material. This accumulation.  48–60 yellowish brown compact clay containing many light grayish tagments of lime 48–50 into estabonish containing many light grayish kayens, the material being partially containing many light grayish kayens, the material being partially connented.	0–8 brown to light brown sandy loam darkened by presence of organic mattler.  8–24 light brown sandy loam. Both this and the above horizon contain little gravel, and considentable coarse sand. Both horizons are loose and friable and contain numerous grass roots.  24–30 grayish brown gravelly sandy loam. Slightly compact	6—16 brownish yellow friable silty clay loam.  Folls brownish yellow smooth, plastic, heavy, moderately tight day motified light kmy.  The mutities are of moderate oxtent and development; and occur in small irregular veins.  The motifies are of moderate oxtent and development; and occur in grand irregular veins.  The soil maderate medium too irregularly shaped lumps, ranging 17-22 are from \$4 to 12\$ inches in diameter.  The control of 10 12\$ inches in diameter.  The detacked comment heaver clay containing a moderate amount of 18, the motifies. In the art frameutial structure forming 22–30 vertexblard legences from \$5 to 15\$ inches in diameter.  Plastic, reproduced the form of the control of the plastic of the control of the plastic of the control
Jenkintown, Pa			Cincinnati, Ohio
Chester Joans	Dublin olay adobe	Everett gravelly sandy loam	Maddox ailt loam
,	10	9	1-

Table 101.—National Bureau of Standards test sites—Continued.

	Depth of speci- mens	Inches 66	52	98	98	54	30
	Тородгарһу	Level			Slight slope	do Practically level.	Fair Almost level
	Internal	Poor	do	Fair	Good	do	Fair
	Description of soil profile	(9-24 black noncaleureous clay loan. Rather friable. Breaks with concoded fracture into pre-size pieces.  22-12 cantereous transition loger with fourgues of both horizons extending into the layer.  42-88 graysh brown heavy clay loan. Light gray when dry—lighty calcurous.  Below 88 prenet material of old lake hid depocits, Graysh brown concretions that are hard horn energy him.	( 0-10 brownish gray silt loam, slightly streaked with reddish brown.  10-16 gray dom streaked reddish brown and mottled yellowish [10-22 transition to the sentiel brown and thrownish yellow. [16-22 transition to the sentiel loam mottled reddish brown. [16-22 bed of gray graved.	Surface—light brown sandy loun	(f. O-12 dark) brown or brown frashle leam.  12-33 reddish brown or brown frashle leam.  12-33 reddish brown or the clary floam. Moderados vennjaet. Constants of the clark of	(The entire profile is a grayish brown friable, loose, micaceous fine saudy foun containing thin layers of material as heavy as loan and as tight as saud. Stongakareous at surface, and only faintly calcareous at 6 feet.  This soil differs from soil 13 in that it does not contain soluble eurbonates in appreciable amount.	O-56 light grayish brown smooth, friable, micaceous very fine saudy bunnish brown very fine sand. 56-62 light grayish brown very fine sand. 67-63 sann as 0-60. 68-72 sann as 0-60. 68-72 sann as 10-60. 69-64 light in the carbonate form, and formerly called habet alkali.
	Location	Pargo, N. Dak.	Sidney, Ohio	:	Baltimore, Md		Bakersfield, Calif
	Soil type	Fargo clay loam Fargo, N. Dak	Genesee silt loam	Gloucester sandy loam Middleboro, Mass	Hagarstown loam Baltimore, Md	Hanford fine sandy loam Los Angeles, Calif	13a Hanford very fine sandy loam. Bakersfield, Calif
l.	Site No.	æ	G	01	11	12	13a

	44	36	30	36	\$
Very gently undalating.	do	•	Gentle slope	do	Practically level.
Good	Fair.	}Poor	Fair	do	(tood
0-6f grayish brown very slightly comparted bunn	0–15 dark brown (almost black) silt loam  Ley transition layer consisting of tougues and streaks of the two adjoining horizons extending into each other. 22–42 brown silt loam with yellowish eask, slightly compact. 42–4 grayish brown sand containing some gravel [Entire profile is nonseleareous.	(0-36 black clay with no appreciable change. Highly calcareous.) Small fragments of line are found throughout the section	0–8 grayish brown fine sandy loam, which appears to have been designatived.  8–42 yellowish brown very fine sandy loam. Texture gradually gets finer and compactness increases with depth. Some reddish muchtings and at few iron concretions about 3 in then in diameter, which are most numerous at about 3 inctanged with red.—which are most numerous at about 3 inctand disappear at 6 fret.—48–48 brownish yellow git loam mottled with red.—48–60 mottled red, gray, and yellow material containing thin layers of elay and fine sand but with the average texture of sile toam. Below 25 inches the color is light yellowish brown with light gray mottlings.	0-6 grayish brown loam or silt loam without structure. Mod- ord, transition and fraible compart clerk four- ord, transition have alightly compart clerk loam.  orded Branch and transition with con- colded fracture sets at the full price with invested clerk.  148-74 frown from sensor free free with invested clerk.  148-74 frown from sandy Joan with slight reddish casts.  151-16 light gray clarky sensor.  151-16 light gray clark gray clark gray sensor.  151-16 light gray clark gray gray gray gray gray gray gray gray	0-8 dark brown silk loam full of brickbats, plaster, rotten wood, the The surface soil partly removed and mixed with foreign matter.  8-72 light brown very uniform smooth fribble silt loam that gets a light brown spot added to the foreign with the silt brown spots due to rotten roots at 8 to 24 inches. Very faintly calcarcous at 48 inches and below.
ор	St. Paul, Minn	San Antonio, Tex	Mobile, Ala	Alsxandria, Va	Omalu, Neb
do	Hempstead silt loam	Houston black clay	Kalmia fine sandy loam	Keyport loam	Knox silt loam
13b	14	12	16	17	18

	44	36	30	36	\$
Very gently undalating.	do		Gentle slope	ор	Practically level.
Good	  Fair	Poor	Fair.	do	Good
0-6 grayish brown very slightly compacted loan.  6-84 light grayish brown friable loose micaceous very fine sauly loam. Namerous rooks in first 3 feet. Few light colored speeks at 3 feet.  A 8 special set of specimens are buried at the site. The profile is similar to site 13a, but differs by being low in ablail content.	0-15 dark brown (almost black) silt loam. 13-24 transition layer consisting of loanes and streaks of the two adjoining horizons extending into each other. 24-42 brown silt loam with yellowish east, slightly compact. 24-45 rayish brown sand containing some gravel. Entire profile is nonealearrows.	f 0-36 black clay with no appreciable change. Highly calcareous. Small fragments of lime are found throughout the section	0–8 grayish brown fine sandy loam, which appears to have been disturbed adjacently and a proper set and compendates inverses with depth. Some reddish mothlings and a few from convertions should fit inch in diameter. Which are nest numerous at bour for set and a few from convertions about fit inch in diameter. 42–48 brownish yellow or yellow slit hoam mothed with red	0-6 grayish brown loam or silt loam without structure. Mode- centely loos and frish sightly compart clay loam in 1-4-4 lightly sublawish bown rather compact clay loam with con- ordal fracture expesing slints surfaces. Slightly mottled with Rey 2 costure gots a little lightle with inversable clash. As 34 brown fine sandy toan with slightly redish cast. The 10 light gray clayey sand. The 10 brown sand almost saturated with water. Batter profile is noneatenrous.	0-8 dark brown silt loam fall of brickbate, plaster, rotten word, etc. The surface soil partly removed and mixed with foreign matter. 8-72 light brown very uniform smooth friable silt loam that gets a little light brown very uniform smooth friable silt loam that gets a little lighter in eader with depath. Moderately moist, Contains a few brown spack due to refer routs at 8 to 24 inches. Very family calcureous at 48 inches and below.
ор	St. Paul, Minn	San Antonio, Tex	Mobile, Ala	Aləxandria, Va	Отаћа, Међ.
qp	Hempstead silt loam	Houston black clay	Kalmin fine sandy loam	Kayport loam	Knox silt loam Omaha, Neb
13b	14	15	16	17	18

Table 101,—National Bureau of Standards test sites—Continued.

	Depth of speci- mens	Inches 36	48	9		90
a new rate of the second	Internal Topography drainage Giord Moderate slope.		Moderate slope. Gently undulating.		Very gently undulating,	Level
			Poor	Good	op	Poor
	Description of soil profile	Description of soil profile  1-4 dark brown silt loam, friable and full of organic matter  4-18 slightly compact heavy silt loam, yellowish brown.  18-34 transition layer into  18-35 rather compact more yellowish brown day containing a few  18-36 rather compact more yellowish brown day containing a few  18-36 gaysish brown day loam with bright yellow mottlings and a  18-36 gaysish brown day loam with bright yellow mottlings and a  18-36 gaysish brown day loam with bright yellow mottlings and a  18-36 gaysish brown day loam with bright yellow containing light  18-36 brough and the standard of years of the standard color, containing light  18-36 brough and the standard brownish gay heavy silt loam or light silty clay loam.		0-28 brown or chocolate brown friable, uniform silt loam. 28-36 themsition layer. 36-38 light brown silt loam very uniform and smooth. Non-calcarcous to 6 feet. 34-light brown noncalcareous clay slightly mottled with grayish brown.	0-4 light brown silt loam containing thin discontinuous layers of darker olor probably due to the turning under of organic matter when the soil was cultivated. 4-60 light brown slightly compact silt loam with some grayish mothlings but no hard lime concretions. Very uniform in color and texture.	0-14 dark brown (almost black) silt loam. Y-inch crust, 3-inch mulch, which is underlaid by slightly compact very lightly moist material with no definite structure.  14-72 light gray loam, moderately compact and moist with somewhat fighter texture and a more open structure below 48 inches, where thin layers of sandy loam occur. Friable and loose.  Thin layers of grayish brown sand occur at 60 inches.  Location has all indications of a soil high in alkali. Highly calcarcous up to surface.
	Location	Des Moines, Iowa	Cleveland, Ohio	Kansas City, Mo	Memphis, Tenn	Buttonwillow, Calif
	Soil type	Lindley silt loam	Mahoning silt loam	Marshall siit loam	Memphis silt loam	Merced silt loam
	Site No.	130	20	21	52	23

¥	36	36-48	<b>2</b>	Ş	12	*	77
:	:		Level	Level to gently rell ing.	:	Level	Almost level
Good	  Fair		Very poor Lavel .	Poor	Very poor	Poor.	Good
0-4 brown loam containing considerable sand and coarse sand)(Gowl 4-33 + grayish coarse sand or fine gravel	O-6 grayiah brown silt loam. 6-30 yallowsish brown, sift, heavy clay loam to clay, containing a small amount of gridy material. 70 8-8 siltingly quata consistoronish yallow heavy clay loam, somewhat 18 iliding than the above and take contains some grity material.	9–2 grayish brown silt loam. 2–7 forwaish gray of solbreish-gray silt loam. 7–10 gray silt loam nottled fairly with yellow. 7–10 gray silt loam nottled fairly with yellow. 7–2 to gray silt loam to ottled fairly with yellow and gray silt loam. 7–2 to year clay loam to clay nottled brownish yellow and yellow. 24–36 to endlah brown stiff clay. 30–38 oeddah brown stiff clay. 30–38 oeddah prown stiff clay. 30–38 oeddah prown stiff clay. 30–48 oeddah brown stiff clay.	(Dull red heavy calcarsous clay extending down below the depth at which the specimens are buried. Soil map shows Milner clay at this location and a sample of the soil was identified as typiral Miller clay.	0-8 filled material—briokbuts, gravel, etc. 8-46 grav or light gravals-brown adobe containing soun gritty material and gravel in the first foot. 46-56 light gray sandy olay, semewhat sticky. 66-60 graylah brown or yellowiah brown gravelly sand. (64- graylah brown or yellowiah brown gravelly sand.	Surface—to varying dopults consists of dark colored material of variable to extern, note of which for a composed mutch which Russed—black, scanfluid mass of well-decomposed mutch which rest upon at allines whill mate of die topress scamps and roots that are in an excellent take of preservation. Substantine—stiff publishe gray and supplied to the servation. The start was originally a cypress swamp.	0-6 dark brown silt loam (grayish brown when dry). 6-72 gray or grayish brown silt loam with yellow mothlines that are wenly distributed and containing a few brown speeks. Non- calcarceous throughout.	0-4 grayish brown five sand containing organic matter
Norwood, Mass	Milwaukee, Wis	Springfield, Ohio	Bunkie, La	San Diego, Calif	New Orleans, La	Davenport, Iowa	Jacksonville, Flu
24 Merrinac gravelly sandy loam. Norwood, Mass	Miami elay loam	Miami ailt loam. (motilied phase).	Miller clay	Montezuma clay adobe	Muck	Muscatine silt leam	Norfolk fine sand
77	% 81 <b>4</b> 786	% ~_4819	72	28	57	30	15.

TABLE 101. -- National Bureau of Standards test sites—Continued.

	Depth of speci- mens	Inches 48			*	<del>9</del>
	Тородгарћу	Gently slop- ing to un- dulating.		Gentle slape.	Moderately rolling.	Gently rolling.
	Internal drainage	Gnod	Very poor	  Good	Fair	Good
The second secon	Description of sail profile	0-8 brown to grayfal brown (when dry) mellow and friable, fine sandy Joan to fine sand. 8-18 shightly more compact, though erunbly bann to fine sandy Joan, lightly more compact, brown in ealor. 18-33 grayfal brown to relievable brown in color. 18-33 grayfal brown to for brownish gray compact learn in place, though friable when lorded out. 18-34 practially weathered (Ill material L	(A black well-decomposed peat 30 to 36 inches deep, where it resis on a drab or bluish plastic clay foam. The lower part of the section was saturated with water. The peat nerges into clyde loam, the line of separation heigr rather indefinite.  A sample of this soil lost 42 percent on ignition.	0-8 brown or dark brown silt loam. 8-24 reddish brown silt loam containing considerable sand. 24-38 slightly lighter in color than above layer. 38-56 Indian red or reddish-brown silt loam. 56 + shale.	0-22 light brown moderately compact loam with slight reddish this and a slight admixture of organic matter to 2 inches of surface. Very dry. 22-5-54 slightly moist, bard, grithy, compact, britds, reddish brown day loam containing numerous white speek. 54-72 light reddish brown or light-brown gritty silt loam. White speeks present but not as compact as horizon above. Entire profile is noreal-arrows.	0-8 light brown, loose, friable sandy loam 8-30 brownish red or rusty brown heavy fine sandy loam. Rather compact and hard. 30-60 reddails brown, rather compact, heavy fine sandy loam. (60-96 mottled red and yellow compact, heavy fine sandy loam (No gravel or stones present in the profile
	Location	Rochester, N. Y	Milwaukee, Wis	Norristown, Pa	I.os Angeles, Calif	Meridian, Miss
	Soil type	Ontario loam	Peat	Penn silt loam	Ramona loam	Ruston sandy loam
1,	Nite No.	25	88	32	35	98

30	30	98	30	8	30
Practically lovel.	Moderate uni- form slope.	Practically lovel.	Gently un- dulating to level,	Gentle dope.	Steep slope
Poor		Fair	Poor	Fair	Pour.
0-2 dark gray or grayish brown fine sand. The organic matter imparts he dark color.  2-10 the material merges into a rather compact yellowish layer baving a distinct lower boundary. The organic matter decreases with depth and the yellow color becomes brighter. The yellow sand contains a few very hard round black iron concretions about ½ inch in diameter that are surrounded by reddish brown sand.  10-28 light exp slightly compact fine sand which becomes lighter with increasing depth and is almost white as 28 inches.  28-36 dark brown hard compact iron cemented hardpan with the characteristic office ground color.	0-8 grayish brown gravelly sandy leam which gradually changes into a lightly vellowish invon or yellowish gray.  8-28 light gray or yellowish brown gravelly sandy leam which is advice that the horizon below.  28-30 light gray gravelly sandy leam with faint yellow east	This soil has been so disturbed that an accurate description of the profile is impossible.  1-12 grayish brown moderately friable silt loam.  0-30 + silghtly yellowsish-brown silt loam which extends below the specimens. The teameh bottom shows considerable gravel and a little gravel exists throughout the profile.	0-8 dark brown or brown elay loam containing organic matter and full of grass rooks. Rather ampact, service as 8-30 stiff, plastic gray clay motified with rusty colored material. No definite mart bro concettlement on concettlement 30-70 gray stift loam motified with rusty brown. The rusty colored spots got girls stift loam motified with depth and practically disappeared at 60 moties.	0-22 very uniform and smooth brown silt loam. 22-36 light brown smooth silt loam. 36-108 light brown uniform silt loam taindy motitied with graysish brown. Nonenleaveous to 9 feet at which depth the soil is underlain by slutte.	Top soil corroded away.  -6. father common but friable light reddish brown chay.  6-45 motified red, yellow, and gray yver what compact that land a calcined structure.  In a calcined structure.  45-56 motified red, yellow and gray heavy silt loan.  56-86 same as 6-45.
Jacksonville, Fla	Camden, N. J	Wilmington, Del	New Orleans, La	Капква Сіґу, Мо	Meridian, Miss
Sk. John's fine sand	Sassafras gravelly sandy loam.	Sassafras silt losm	Sharkey clay	Summit silt losm	Кияспећапла. свау
37	88		40	14	42

Table 101. -- National Bureau of Standards lest sites—Continued.

Batire soil profile, and especially the surface foot, contains a large percentage of undecayed organic matter and has a black color when wet. Upon drying the color clinages to grayish brown. The soil contains hydragen suffice and a considerable amount of soluble sails, but no line. The surface portion of the soil lost 20.7 percent on ignition.
Except for the addition of gruss roots to the top 8 to 12 inches, the entire profile consists of a uniform dark brown silt loam (thlack when web or silty chay loam, to a depth of at least 8 feet. Non-calcarcous throughout.
0-6 light gray to light grayish brown sand to heavy silt loam.  Jittle organie matter.  2-20 brown to grayish brown heavy, compact, gritty clay.  Plaste and waxy when wet, but becomes lard and tough when dry.  20-20 abrupt clanage to a light gray sandy clay. More friable than upper horizon due to light gray sandy clay. More friable off an organic decreases, color slightly darker and texture more compact than above horizon.  Type is highly alkaline, and white streaks and subotches of con- centrated salts occur abundantly throughout the profile except in the surface soil.
14-20 light brown or light brown sandy loam. 14-20 light brown sand debris 14-20 light brown sand debris and friable. 20-22 lard compared layer of cinders. All the above material is lose 20-22 lard compared layer of cinders. All the above material is profile. 22-36 lard. compact brown sandy loam. 22-36 lard is lightly enemerated.
0-12 grayish brown or brown silt loam containing considerable organic marter. Highly etlacuous at hall dight, so. 12-72 light gray moderately compact clay containing consional mothings of brownin, yellow and reddish brown. A few lime concertions and occasional water-worn pebbles that are partly coated with lime are present.

90	98		39		30		:	95
Level	ор		Steep slope		do			do
Very poor	do		Poor		do	do	Very poor	Роиг
The area is a transition from Acadia clay to prarie of Lake Charles clay. The telestive in in the two soil types. The 20 feet of bouth end of trench is Lake Charles clay. Acadia clay, prairie phase. 0–12 very dark gray (almost black) heavy acid clay spotted with yellowish brown.  12-30 dense gummy dark-acid clay with yellowish brown and rust brown spots and splotches.  Large amount of line soil crystals of gypsum, neutral in reaction. Lake Charles clay.  0-2-4 black heavy clay.  2-4-40+ yellow heavy clay with some gray mottling and fine crystals of gypsum.	0–12 dark gray silt loam. White incrustation of soluble salts on the surface of gray gilty clay loam nottled with yellowish brown, containing some black concertedions.  20–20 + gray and yellow dense gummy mottled day containing a few addition arthonic conrections. Pharent material of eal-eaceous elay lies several feet beneath the surface.	Same as site 3.	0-5 gray or light yellowish gray gritty, friable, silt loam stained or speeded with light gray and rush brown. Moderate quantity of small calearcous shale chips present. 6-12 light gray or light brownish gray gritty, slightly compact friable silt loam constaining a largen amount of small chips of calearcous shale and linestone. 12-24 gray calearcous thin beds of shale partly weathered to clay stained light gray, alloarcous shale containing small irregular pockets of gray, plastic, leavy only or partly weathered shale.	Same as site 11.	0-12 black noncatearcous, very heavy clay.  12-33 dark bluish gray, noncatearcous, waxy clay.  32-44+ light gray waxy, noncatearcous, clay with some yellow sputs.	Same as site 117.	Description not available. Soil very similar to site 29.	0-13 black or vary dark gray granular, smooth, bomy, thoroughly decomposed organis material. Anderartely and control of 13-30 dark gray fibrous, stringy, moderately compact plant remains partially decomposed and containing brown raw felly post. Slightly and, partially decomposed remains on swamp-byring plants displayed.
Spindletop, Tex	League City, Tex	Atlanta, Ga	Cincinnati, Ohio	Baltimore, Md	El Vista, Tex	Tranquillity, Calif	New Orleans, La	Kulamazao, Mich
Acadia olay	Lake Charles clay loam (mound phase).	Cecil clay loam	Fairmount silt loam	Hagerstown loam	Lake Charles clay	Merced clay adobe	Muck	Curlisle muck Kulamazao, Mich
51	29	23	54	22	56	57	82	99

Table 101,--National Bureau of Standards test sites—Continued.

Depth of speci- mens	Inches 22				24-30
Тородгарну	steep stope			do	Level
Internal	Very poor			ob	Very poor
Description of soil profile	0-4 dark brown or brown loanny, smooth well-decomposed organic matter internived with finely dibrous, partly decayed vegetable matter that includes some woody undertal in different stages of denomposed organic matter that includes some woody undertal in different stages of the control of th	Same as site 40.	Same as site 42,	(This soil is typical of the titlal marshes found along the seacoast. The soil materials from titlal marshes vary from dark, coay sediments interspersed with coarse marsh-grass roots to a yellowsh or dark colored lady. The subsoil and the free water in the soil are usually charged with hydrogen sulfide.  A similar soil is singed with hydrogen sulfide.	O-12 very dark gravish brown clay that is plastic when wet and has fairly lurge enclass when dry. Soil blooky and moderately compact when dry. Many salt crystals.  12-24 gravish brown day that is plastic when wet and has frable consistance when moist. It has less visible lime than the surface soil but is highly calcarcour. As seen that it contains many lime splotches.  24-36 similar to layer above, except that it contains many lime splotches.
Location	Plymouth, Olifo	New Orleans, La	Meridian, Miss	('harleston, S. C	Cholame, Calif
Soil type	Rifle peat	Sharkey clay	Susquehanna clay	Tidal marsh	Docas elay.
Nite No.	8		62		79

98	34-48	88
Almost level	do.	J.ovel
do		Very poor
This soil is a poorly drained phase of Chino silt loam.  0-13 dark-hoven friable, indexons, highly addarcous silt loam motified with rust brown and gray silt loam or elsy loam.  13-28 hownish gray highly motified with rust specks, firm elsy loam that is highly achaerous.  23-30 highly achaerous olive gray and grayish brown motified, very fine sandy loam, not so much iron motifiing as layer above, and becomes light gray when first load pooms wery fine sundy on the sand to an and becomes light gray when fury.	This area is mapped as Mohave fine gravelly loam (lighty calcareous phases), but it would now be classified as Thoson fine gravelly loam. Within the length of the trench is a thin stringer of Pinh fine gravelly loam. These two solis grade into each other and are therefore not typical of either. Descriptions of the two soil profiles (1010 v	Tast site is adjacent to a canal and the treach is always more or fless we. The treach as evenemed to a cloud of 61 feets and filled with fresh cinders, in which the specimens were buried to stee, An analysis of the cinders showed it. Ve contain 20 percent of carbon. Following is the description of the profile before hackfilling with the cinders, and carbon. So and ye day with article of a mixture of cinders, canal dust, and oxide.  8-17 layer great and shavings with about 60-percent sulfur content and some granide compounds.  17-60 mixture of sandy chay and loam with appreciable quantities of mark and traces of inders and ashes.
Wilmington, Calif	Phoenix, Ariz	Milwaukee, Wis
Chino silt loam	Mohave fine gravelly loam. (Tueson fine gravelly loam).	Cinders
65	99	29

TABLE 101. -- National Bureau of Mandards test sites - Continued.

	Depth of speci- mens	Inches 40			ope.
	Тородгарһу	Level	do		Gradual slope.
	Internal drainage	Imperiest	.Poor		Very poor
to the supplier of the suppline of the supplier of the supplier of the supplier of the supplin	Description of sail profile	0-12 uniform chocolate brown granular, highly calcarcous chay that is frable when moist, This layer is quite plastic when moist brit not so much as the layers below.  The Audion brown massive, firm, highly calcarcous clay which grades into.  24-38 more friable red tinted brown, highly calcarcous clay that languages into.  36-88 sightly models red, inted brown, highly calcarcous clay that sightly models red, as well as highly calcarcous clay that is every plastic when wet.  48-90 + very little change in texture or hime content. Motthings	0–8 vory dark gray or dark grayish brown smooth, loanny, thoroughly decomposed organic matter matted together with plant roots and interminged with raw or partly distingated finely fibrous peatly matter. Contains some sand material. 8–24 dark gray smooth, listle, nouny well-decomposed vegetable matter containing finely divided partly decayed plant remains and brown raw fibrous peat. Layer thinly laminated and indefentiely compact and matter with fine roots penetrating through it.  24–35 brown raw, fine, fibrous peat containing dark or gany smooth, loanny partly decayed organic matter and matter and matter and matter with such years when yearly decayed organic matter and matter and mater and matter and water and mater and water and material.	Same as soil 23.  Description of profile not available. Soil type is subject to change upon better identification.	This soil is a dark bluish black when wet and has a pH of 7.2. When art-dried it is hight gray with a brownish east and has a pH of 4.1. Active anaerobic sulfide-reducing bacteria have been identified here. Argued profile from soil survey report of Portage County, Surface—varies from a clay fount to a silty day loan, generally of a somewhat bluish east, rvey wet places. The dirty profiton have a brownish east. The soil, while heavy, earries sufficient, sand to be noticeable. Subsoil—consists of a clay loan to silty day loan, light gray or gray mottled with some drub now. Day content increases with depth and mottling usually becomes or proven a steel-blue, soft, plastic city. Presence of iron stains or your a steel-blue, soft, plastic city. Presence of iron stains
	Location	Phoenix, Aria		Buttonwillow, Calif Austintown Junction, Ohio.	Yale, Ohio
A STATE OF THE PARTY OF T	Soil type	Gila etay	Houghton muck	Merced silt loam	Trumbull clay loam
11	Site No.			2 2	22

						# 	8	98	<b></b>	93	¥ .	75
						Very gentle uniform slope.	Very gentle slope.	:	Gentle slope.		do	do
					-	Good.	Fair	do.	Good	do	du	
Description of profile not available. Soil type is subject to change upon better identification.	do	op	do	op	do	0-6 gray silt boan containing a few cinders and showing evidence of brain gisturior. Rather compact and dry. 6-15 graysh brown slightly compact silt foam full of white streaks and spots. 45-108 grayish brown sandy loam, loose and rather wet at 108 infelies.	0-6 grayish brown silt loam, lighter than lower soil probably beause It is dryv e-108 grayish brown silt loam mottled with white streaks and species and containing a few thin layers of fine sandy foam below 58 meles.	0-2 surface erust underlain by mulch, grayish brown siit loam 2-74 grayish brown heavy slif Joann. White erestka and speaks appear below 14 inclets and become most numerous at about 74-108 light grayish brown micaecous fine sandy loam.	0-72 bright red clay, smooth and brittle with no noticeably change, except slight addition of organic matter in first few inches.	0-8 grayish brown sandy chay loan, pine needles and trash, above 1-8-28 red brittle chay commert and hard 28-73 minesons little tot did bush to fall the 1-72-100 methed red and yellow minescons with loan. No visible granibe, but so have account of the period of the structure-like petrified word.	(0-4 grayish brown clay loan. 4-72 red chy which gots a little more crumbly and slick with increasing depth.	0-8 grayish brown fino sandy loan with a little gravel mixed with it, taker compact. 18.1 stychowsh frown chay hean, more frable. 18.3 tree lay. 18.4 tree lay. 18.4 tree lay. 18.4 troubled, grilly red and yellow partially decomposed granite.
	Rocky Ford, Colo	Albuquerque, N. M	Los Angeles, Calif	Louisville, Miss	Latex, La	Grand Junction, Colo	Grand Junction, Colo	op	Charlotte, N. C	Macon, (ia	Salisbury, N. C	Raleigh, N. C
73 Unidentified river bed E. St. Louis, Mo	Otero clay loam	Unidentified alkali soil	Chino silt loam	Susquehanna clay Louisville, Miss	Caddo fine sandy loam Latex, La	Billings silt loam (low alkali).	Billings silt loam (moderate alkali).	Billings silt loam (high alkali) .	Cecil clay	Cecil clay loam	ф	Cecil fine sandy loam
73 U	74 Ot	75 U	76 CI	77 Su	ت 28	101 B	102 I	103	104	105	106	107

Table 101,--National Bureau of Standards test sites-Continued.

	Depth of speci- mens	Inches 36	36			98	36		
	Topography	Gentle slope.	. Lovel	do		Gentle uni- form slope.	do		Gentle slope.
	Internal drainage	Excellent	Poor	qp	do	do	Fair.		Good
	Description of soil profile	0–2 grayish brown sand loam 2–42 typical red city 42–72 red city with mica crystals	O-2 loose grayish brown fine sandy loam.  10-18 by hard, comparing grayish brown, microcous fine sandy loam containing a few light colored species. Breaks with correcting fracture.  18-64 grayish brown compact shale-like microcous very fine sandy four containing rusky mottling in places. Lighti gray streaks between plates.	O-6 gray moderately compact fine sandy loam. 18-21 compact light bown fine sandy floam. 18-23 way hard, compact, tough layer, breaks with conocial fractum. 24-86 moderately compact light brown ailt loam. 18-68 light brown moderately compact fine sand. (18-72 sane as above with rusty mottling.	Description of profile not available. (See site numbers 109 and 110.)	0-50 brown shale-like compact clay with thin layers of a coarser material and gravel scattered through the profile. Practically no gravel on surface. 50-56 layer containing considerable gravel. 56-72 brown clay.	1-4 light brown erust. 1-4 light brown midd. 4-80 brown glay with a slightly reddish or purplish east, hard and compact with lanky structure. If the entire profile had the same moisture content, it would probably have the same color. Texture is heavy throughout and practically no gravel or coarse material.	Same as soil 56.	(0-12 grayish brown silt loam. 12-72 light brown silt loam with slightly reddish cast. No lime or mottling.
***************************************	Location	Atlanta, Cia	Presno, Calif	ор	Kernell, Calif	Niland, Calif	do	El Vista, Tex	Vicksburg, Miss
	Soil type	Cecil gravelly loam Atlanta, (fa	Fresno fine sandy loam (low Fresno, Calif., alkali).	Fresno fine sandy loam (moderate alkali).	Fresno fine sandy loam (high alkali).	Imperial clay (moderate alkali).	Imperial clay (high alkali)	Lake Charles clay	Memphis alt loam.
il.	Site No.	108	100	110	Ξ	112	113	114	115

	24	30		0#	36			
	Level	Moderate slope.		Level	ф.	:	Хеер вюрея.	
Fair	Poor	Fair	Good	·····op· ··{	do	Fair	Poor	do
Description of profile not available. Profile most similar to site 117.   Fair	0-3 loose grayish brown mulch, showing characteristic adobe 7-18 dry, grayish brown elay, showing characteristic adobe eracks. The lumps are very hard, and the cracks are lined with light colored salts. Profile supears as though there was a heavy concentration of alkali at shout 18 inners.  18+ moist elay with no well-defined structure.	(0-26 stratified light gray or grayish brown gravelly sandy beam. 28-28 brown compact clay. 32-38 brown compact clay. 32-48 light gray gravelly sandy loam—same as surface clay. 54-72 light grayish brown sandy clay.	0-4 graylah brown sandy loan. 4-60 yallowish brown or dirty yellow fine sandy loan. These two layers are nulter consistency of the party layer of variable thickness.	$\left\{\begin{array}{lll} 0-2 & \text{yellowish gray sandy loan.} \\ 2-72 & \text{pale yellow fine sand with some elay mixed with it.} \end{array}\right.$	0-4 gray or grayish brown.  4-12 gradually changes into pale yellow sand. Color typical of Norfolk at 12 inches.  112-65 yellow color increases in brightness and deep thu util it is a slight brown with brownish yellow not	Typical profile from reconnaisastone soil nurvey of the lower San footnin Valley, Chlif. Pancelee deel Jonnia is gray of provinish gray that interconnection. Subsolit occurs at a depth of 18-30 nations and resembles the surface in essential issuers. Call arreous, derevening freigh with shall, and the limit in many places is concentrated in seams, veins and notities. The deepper subsoli and notities and obstactions includes strata of small gravel, and, and finer sediments.	(4.2 reddish brown fine loose sandy loam. 2.4 red compact take for red (5.2 yellow elay. 14-25 red compact and red (5.3 yellow elay. 42-25 yellowish gray untilled with light gray elay. 12-72 busin gray compact elay.	Typical profile from soil survey of Rapide Parish, J.a.  4-1 brands in Grayhai sill loam.  4-12 pallo yellow or mottled yellowish and grayish silt loam.  12-1 yellow clay, which becomes mottled first with gray and in the lower part with red and drab. Jower subsaul is very plastic and adheeve when wet.
Los Banos, Calif	Tranquillity, Calif	Niland, Calif	Macon, Ga	Pensacola, Fla	Tampa, Fla.	Mendota, Calif	Shreveport, La	Troup, Tex
116   Merced clay	Merced clay loam adobe	Niland gravelly sand (low alkali).	Norfolk sandy loam	Norfolk sand	op	Panoche clay loam	Susquehanna olay	Susquehanna silt loanı
116	117	118	119	120	121	122	123	124

Pable 101,—National Burcau of Standards test sites—Contine

	Depth of speci-		_	obe.			<u> </u>
	Topography	Moderate slope,		Clentle slope.	Lovel.		do
	Internal	Poor		Poor	Good		Very poor
the contract of the state of the states of the state of t	Description of soil profile	(P-10 light-hrown slightly commet fine sendy leam.  10-30 yellowish-hrown rather compact elay lean containing some 30-48 mottled real, yellow, and gray silt leam.  48-72 mottled real, yellow, and dray silt leam.	AMERICAN PITROLEUM INSTITUTE SITES	Typical profile from soil survey report of Bell County, Tex. 12-30 gray, heavy plassic clay. 30 + gray clay.	Soil described as black, tough, heavy clay. Location of the site on the less of map accompanying the soil survey of Cowiey County Kans, shows the soil to the Oswegistil toun. The typined profile by a clay hardpen or onesisting of black, tough, heavy clay in material shows little change within the 3-local section, except that in the lower part it is frequently lighter colored heing very dark fanb to very clark known or marry black with some yellow is the black capture concerning and the colored concertionary material. This type is the black captured to Gerald slift beam, it is locally known as "heady and fair feather of Gerald slift beam, it is locally known.	Same as National Bureau of Standards site 56,	This test site is in the same are as site 22 of the National Bureau of Shandards site. The Makional Bureau of Shandards site. The Makional Bureau of Shandards specimens were buried in the mound phase, which is different from the remounding soil in which these specimens were buried. Pripatel profile from soil-survey report to Galveston County, Tex. 124 gray chay.
37	Loration	Streveport, Lat.	AM	Temple, Tex	Arkansas City, Kans	Beaumont, Tex	
The state of the s	Soil type	Susquehanna fino sandy loan.	-	Bell clay Temple, Tex	Oswego silt loam	Lake Charles elay Beaumont, Tex	Lake Charles olay loam League City, Tex
	Site No.	125		-	61	. S	4 I

							<del></del>		
	op	lo	ор.		do	ор	do		
	:	do	:		:	:	:		
	Poor	}do	do		Good	Poor	}do:		_
Typical profile from soil-survey report of Blackford County, Ind. 0-2 dark gray loose silt loam mixed with well-decomposed organic	2.6 pale yellowish, gray frable silt loam that is soft and crumbly. The lower half contains vey littled regains matter. 6-12 heavier silt loam, which is rather firm but porous and has little or no definite structural arrangement, 124 moderately hard silty elsy loam which breaks into aggregate from 34 inch to 34 inch in diameter. The granules separate easily when moist. Rust brown colorations in the lower part.  34 4 done and lighter-colored massive till, yellow stains occur along the fissures.	Sarface=dark brown friable clay loam  Sabsoil=yellowish brown clay	Sell type amond from location on eals arrays map of Montgomery County, Kans, Typical profile follows: County, Kans, Typical profile follows: County, Gark was been a sell follows: Self black gray to black sill know malterlain absurptly. Self loca not chalmed with depth, but the color usually becomes self there are the maps with depth, but the color usually becomes self there are the proven color. The sail at this test sich has been inditified as follows: Surface—black silt foam. Subsoil—lieuvy yethowsh foam.	Same as prairie phase at National Bureau of Standards site 51.	Soil type named from location on soil-survey map of the Anaheim Area, Calif. See National Bureau of Standards site 12 for typical profile.	Typical profile from scil-survey report of Promediate County, Iowa. 10-2 vor dark grapish brown or havy black till hann. 2-18 vorg dark grapish brown or havy black till hann. 2-18 vorg dark grapish brown sil hann that is very viciely vicin wet but frished welled from the the series of the scill be levely in texture, anymorabing pally day from John John 18-25 volled with frown tilly day John	(Upper soil—striated brown sandy clay. Subsoil—black clay. Most of the line is in a salt wasl.	Same as National Bureau of Standards site 117.	13 Miller clay Bunkic, J.a   Bunkic, J.a   Same as National Bureau of Standards site 27.
	Preble, Ind	Council Hill, Okla	Cancy, Kans	Spindletop Gully, Tex	Long Beach, Calif	Mt. Auburn, Ill	Skiatook, Okla	Mendota (Tranquillity), Calif.	Bunkie, La
	Miami silt loam	Unidentified	Oswego silt loam	Acadia clay	Hanford fine sandy loam Long Beach, Calif	Muscatine silt loam	Unidentified	Merced clay loam	Miller clay
_	10	9	7		6	10	11	13	13

Table 101.—National Bureau of Standards test sites—Continued.

Depth of speci- mens	Inches		98		21	18-24	¥2	- 24
Тородгарћу	Gentle slope.		Level		Level	Very gentle uniform slope.	Depression	do Level
Internal	Good		Good		Very poor	}do	do	op
Description of soil profile	( 0-6 loose, mellow dark brown silt loam.  12.23, stools, yddowith brown silt loam.  12.24 stools, yddowith brown silt loam.  12.24 stools yddown on passe silty clay loam, becoming heavier with dopdis. I dinestone was encountered at various doptis from 24 inches to 36 inches.  Name as National Bureau of Standards site 04.	AMERICAN GAS ASSOCIATION SITES	The soil is all fill (at least 10 years old) to a depth considerably below the bottom of the trench. It consists mostly of cinders from the gas plant, but also present was some bolin-fule dust, as from reforts, only, concrete, a little gravel, and other material. The dark colored and eleaner cinders were placed next to the specimens.	Same as National Bureau of Standards site 67,	This site is covered with about 6 inches of water at high tide.  The soil consists of 12 inches of a thick mat of grass roots, which is underlain by gravel and sand ninted with organic matter. The lower part of the first foot is darker colored than the surface 6 inches, and it has a distinct odor of hydrogen sulfide.	( 0-24 heavy mat of grass roots. (24 + loose, semifuid mass.	Muck is above sand, being about 3 feet deep at one end of the trench and a few inches deep at the other end. The sand was thrown out of the trench and muck placed next to the specimens to a depth of at least I foot all along the trench. Most of the muck is black, but it contains a few lenses of brown fibrous peat.	(0-8 black well-decomposed muck 8-16 transition layer. 16-56 brown fibrous peat. 66-72 black semilluid sand, which is white sand containing organic matter.
Location	Chambersburg, PaCholame, Calif,	-1	Pittsburgh, Pa	Milwaukee, Wis	Brockton, Mass	Atlantic City, N. J	West Palm Beach, Fla	Miami, Fla
Soil type	Hagerstown loam		Cinders Pitisburgh, Pa	do	Tidal marsh	do	Muck	op
Site No.	14		-	67	60	4	10	9

8

~	Cecil clay loam	Atlanta, Ga	7   Cecil clay loam   Atlanta, Ga   Same as National Bureau of Standards test site 3.		
00	8do	Raleigh, N. C	0-4 grayish brown sandy loam. 4-8 yellowish brown sandy loam of heavier texture than above hortzon which changes to red lay loam. 8-364-4red lany loam or clay merges into rotten rock characteristic of cell softs.	Fair	Fair Rolling
6	Susquehanna clay	Shreveport, La	9 Susquehanna clay Shreveport, La Same as National Bureau of Standards site 123,		
10	Miller clay	Shreveport, La	10 Miller day Shreveport, La 6-72 dull red or reddish-hrww clay containing occasional thindo do practically level.	do	Practically level.
Ħ	do	Bryan, Tex	11dodo	Poor	do
21	White alkali soil	Los Angles, Calif	12 White alkali sofi Los Angles, Calif This soil is a Hanford fine sandy learn containing a large amount of sufface. Otherwise the profile is similar to the National Bureau of Shandards site 12.		
13	13 Black alkali soil Los Angeles, Calif	Los Angeles, Calif	This soil is a Hanford fine sandy loan unusually high in earbonates. Otherwise the profile is similar to the National Bureau of Standards site 12.		
14	Marshall silt loam	Kansas City, Mo	14 Marshall slit loam Kanssa City, Mo. (20-24 training layer (29-24 training layer (24-1ght bown silt boam)] Excelent Almost level	Excellent	Almost level

ଛ

24 23

# XVIII. APPENDIX 3. METHODS EMPLOYED IN CLEANING OF THE NATIONAL BUREAU OF STANDARD SOIL-CORROSION SPECIMENS

After the soil-corrosion specimens were removed from the trench the loose dirt was scraped off and the specimens were boxed and returned to the National Bureau of Standards. Precautions were taken in packing the specimens to prevent injury to the fragile materials during shipment. When the specimens arrived at the laboratory they were properly identified. The identification letter to identify the material and the number that associates the specimen with the soil to which it had been exposed were stamped on the specimens with steel dies. Each type of material was then subjected to an appropriate chemical and mechanical treatment to remove the corrosion products, with an insignificant loss in weight of uncorroded metal, and to prevent mechanical injury to the specimens.

#### 1. CLEANING OF IRON AND STEEL SPECIMENS

The iron and steel specimens as they came from the field were lightly pounded with a small dull-pointed hammer to remove the lightly adhering corrosion products. If the specimens were in the form of pipes, the caps were removed and the inside of the pipes were washed in a cleaning fluid to remove the coating of grease that was applied prior to burial to prevent internal corrosion. The specimens, which were in the form of plates or sheet, usually had a bituminous coating applied to the two ends to protect the identification numbers. This coating was removed by means of an appropriate solvent.

Up to 1928, the iron and steel specimens were cleaned by pounding them with a small pointed hammer, then brushed with a stiff wire brush, and treated with a bath containing a solution of alkaline ammonium citrate heated to about 80° C. In 1928, air-driven tools were substituted for the hand-cleaning operations and were subsequently used on the iron and steel specimens, with the exception of the

high-alloy steels.

The specimens were pounded with an air hammer employing tools of different shapes and sizes to remove the corrosion products and hard flakes of rust that adhered to the specimens. The specimens were next brushed with a stiff, circular wire brush attached to the shaft of a motor. Then the specimens were placed in an electrically heated enameled iron tank containing a 10-percent solution of ammonium citrate made alkaline by the addition of ammonium hydroxide. The bath was heated to about 80° C. The time the specimens were left in the bath varied, depending upon the amount of rust and the condition of the solution, a fresh solution cleaning them more quickly than a solution that had been used for some time. Usually, immersion in the bath for 2 to 8 hours was sufficient to clean most specimens. After removal from the bath, the specimens were scrubbed under running water with a stiff wire brush and dried with cotton cloths. Frequently, hard flakes of rust still remained on the specimens. These were pounded again with the air hammer and the process repeated until all the corrosion products were removed.

It was deemed advisable to make tests showing the effect of the various tools and of the citrate bath used on the loss of weight of the pipe. Unburied pieces of 3-inch steel pipe and 3-inch cast-iron pipes were used for these tests. The specimens were thoroughly cleaned by the process described. The air hammer was turned on full. For the actual cleaning of the specimens, however, the air is very seldom turned on more than one-third; and only about one-third as much pressure is applied

to the hammer as was applied during this test.

The test procedure was as follows. The specimens were first carefully weighed. One-third of the outside area was hammered vigorously with a sharp-pointed tool; another third with a six-pointed semiblunt tool; and the final third was a sharp-edged tool. These three tools comprise all the tools used in cleaning the specimens. The specimens were then placed under the motor-driven wire brush, after which they were immersed in the citrate bath at 80° C for 8 hours. After removal from the bath, the specimens were scrubbed under running water with a wire brush, dried, and reweighed the following day. The results obtained from the tests are given in table 102.

The maximum total loss of metal due to the cleaning on the 3-inch specimens was 55 milligrams, and on the 6-inch specimens the maximum loss was 320 milligrams. If a corrosion loss of 100 grams is assumed on a specimen, the maximum loss would result in 0.06 of 1 percent for the 3-inch specimens, and 0.32 percent for the 6-inch specimens, an error which is unquestionably negligible for the steel and very slight for the cast-iron materials. In 1937 the greatest accuracy attainable on the balance for weighing 6-inch specimens was 100 milligrams. Also the hammering to which the test specimens were subjected was many times more severe than applied in the

regular cleaning process. The sharp-edged tool, which apparently was the cause of most of the loss due to the hammering, is very seldom used and then very carefully.

This method of cleaning was used for all the uncoated ferrous materials with the exception of the high-alloy-steel materials. These specimens usually did not require any further treatment after scrubbing them with a stiff wire brush under running water. Sometimes it was necessary to remove the corrosion products from the pits with a knife and to place the specimen in the alkaline citrate bath for 1 to 3 hours.

Table 102.—Effect	t of mechanical	cleaning of	f the specimens.
-------------------	-----------------	-------------	------------------

Specimen	Original weight	Weight after using air hammer and wire brush	Weight after immersion in in bath and brushing	Total loss
3-in. wrought iron 3-in. open-hearth iron 3-in. Bessemer steeldo 6-in. cast irondodo	1751.800 1742.660 1765.875	9 1660.440 1751.785 1742.650 1765.870 6880.09 7053.31 6984.70	9 1660.415 1751.745 1742.605 1765.830 6879.92 7053.11 6984.54	0.035 .055 .055 .045 .23 .29

# 2. CLEANING OF COPPER, COPPER-ALLOY, AND ALUMINUM SPECIMENS

The copper, copper alloys, and aluminum specimens as they came from the field were scrubbed with a wire brush under running water to remove the adhering soil. After the soil was removed, all of these specimens were cleaned by placing them in a solution of 5-percent nitric acid and 2½-percent oxalic acid for 5 minutes or less. After removal from the acid bath, the specimens were scrubbed with a wire brush under running water. If necessary, the treatment was repeated. One such treatment was usually enough to thoroughly clean all the specimens except those that had been exposed to Tidal Marsh. The specimens of brass, bronze, and copper from this soil were covered with a very adherent coating of black graphite-like material, which could not be removed or loosened by any reagent so far tried. The wire brush slightly scratches the softer metals but continued hard scrubbing with it on copper and aluminum did not produce a loss which could be observed.

Specimens which had already been cleaned were given the treatment described above. The results obtained are given in table 103. 5 and 10-percent solutions of sulfuric acid were also tried as cleaning solutions for the copper and copper-alloy specimens, but they did not clean as quickly or as well as the solution of nitric and oxalic acids.

Table 103.—Effect of oxalic acid cleaning solution on nonferrous specimens.

		Loss of weight after—				
Material	Original weight	5 minutes in bath	5 minutes more in bath	5 minutes more in bath		
Copper, H. Brass, B. Bronze, L. Aluminum, C1 Al+1½ percent of Mn, C2. Duralumin, C3.	25.68 28.18	0.03 .02 .02 .02 .02 .01	0.00 .02 .02 .01 .00	0.00 .00 .01 .00 .01		

### 3. CLEANING OF LEAD AND LEAD-COATED SPECIMENS

The lead-coated pipes and the lead sheaths removed from the test sites previous to 1937 were scrubbed under water with a fiber brush to remove the soil. They were then placed in a solution of 5-percent nitric and 2½-percent oxalic acids for about 5 minutes. After removal from the bath, the specimens were scrubbed under running water with fiber brushes and dried. If all the corrosion products were not removed, the treatment was repeated. Where the lead coating had failed on the pipe the rust flakes were chipped off with a pointed knife. Ammonium citrate solution cannot be used to remove this rust as the citrate attacks the lead.

Clean specimens of chemical and antimonial lead were given the above series of treatments, with the observed losses shown in table 104.

Table 104.—Effect of	nitric and	oxalic acids	on lead	specimens.

Material	Original weight	Loss of weight after—		
		5 minutes in bath	5 minutes more in bath	5 minutes more in bath
Chemical lead, AAntimonial lead, H	2131.87 2075.04	0.18 .11	0.14 .09	0.02 .05

After 1937 the lead specimens were cleaned by placing them in a water bath maintained at 80° to 90° C for several hours. The specimens were then scrubbed under running water with a fiber brush. The corrosion products were removed from the pits with a dull-pointed knife and again scrubbed under running water. Usually two or three such treatments were sufficient to clean the specimens.

#### 4. CLEANING ZINC AND GALVANIZED SPECIMENS

Up to 1939 the best method that had been found for cleaning zinc and galvanized specimens was concentrated ammonium hydroxide. Dilute solutions of ammonia attacks the corrosion products less and the zinc more than the concentrated solutions. The specimens were scrubbed with a wire brush. Trials showed that a wire brush does not remove weighable amount of zinc even though the specimens were scrubbed harder than is necessary to clean them. The results reported in table 105 were obtained by placing specimens of clean zinc in concentrated ammonia for 5 minutes, which is sufficient time to remove the corrosion products off of most specimens.

Table 105 .- Effect of 10 percent ammonia solution on zinc specimens.

	Original weight	Loss in weight after—		
Material		5 minutes in ammonia	5 minutes more in ammonia	5 minutes more in ammonia
Standard zinc sheet, P Zinc sheet, Z1 Zinc plate, Z2	9 87.92 90.79 504.19	0.04 .07 .06	0.26 .10 .07	0.01 .05 .03

The objections to working with an open bath of concentrated ammonium hydroxide resulted in the adoption of the following procedure for cleaning the zine materials after 1939. The specimens were immersed in a 10- to 15-percent solution of ammonium chloride maintained at 75° to 85° C for 30 minutes. After removal from the bath, the specimens were scrubbed under running water with a wire brush. If any corrosion products remained on the specimens, the process was repeated. It was often necessary to loosen the flaky corrosion products by scratching the surface of the zinc with a dull knife.

Table 106 reports the loss in weight of two specimens of zinc plate.

Table 106.—Loss of weight of zinc specimens caused by ammonium chloride bath and scrubbing.

		Loss in weight after—		
Material	Original weight	30 minutes in bath at 75°C	Vigorous scrub- bing with wire brush	30 minutes more in bath at 75°C
Rolled zinc, Z Die-casting zinc, CZ	509.76 504.00	0.04 .01	0.00 .01	0.04 .03

#### 5. CLEANING OF CALORIZED PIPE

No satisfactory solution has been found for cleaning calorized pipe. Most of the corrosion products are rather loose and flaky. The specimens were cleaned by scraping and brushing the pipe.

### CLEANING OF NONMETALLIC-COATED SPECIMENS

The nonmetallic-coated specimens were scrubbed under running water with a fiber brush until clean and then dried with cloths. After recording the condition of the coating one-half of the coating was removed from the specimens in order to observe the condition of the metal under the coating. The corrosion products were removed from the pits by means of a pointed knife.

### XIX. APPENDIX 4. ACCUMULATION OF DATA, CALCU-LATIONS, AND METHODS OF REPORTING THE DATA

#### 1. ACCUMULATION OF DATA

After cleaning the specimens, the loss of weight of each was determined and checked, with tolerances ranging from 1 milligram (0.000035 ounce) for light stainless-steel specimens to 0.1 gram for cast-iron specimens weighing in the neighborhood of 6 kilograms (13.2 pounds). These tolerances do not, of course, indicate the precision of the data, as corrosion losses were only small percentages of the total weights and ranged from less than a milligram to a few hundred grams. However, the factor that controls the accuracy of the data is the extent to which conditions in the field can be reproduced, and this factor varies within wide limits. Moreover, it is doubtful that this factor can be greatly improved without getting away from conditions to which pipe lines are subjected. Data on the corrosion of working pipe lines indicate that the results of corrosion can not be reproduced within very narrow limits.

After weighing the specimens, the deepest pits were measured and the measurements checked, usually with a tolerance of 0.004 inches. In measuring the pits of the first removals in 1924, the practice was to measure no pit that was less than 10 mils in depth and to measure at least the 5 deepest pits that were over 10 mils in depth. If the specimen did not have 5 pits greater than 10 mils in depth, the depth of the unmeasured pits was assumed to be 5 mils. This assumption was based on the fact that there were unmeasured pits ranging in depth between 0 and 10 mils. Thus, it is fair to assume an average depth of 5 mils.

For the removals made during 1926 and 1928 (second and third periods) pits were not measured when they were less than 20 mils in depth. Thus, on the assumption previously stated, an average depth of 10 mils was assumed as the value to use for

each unmeasured pit.

For the fourth period (1930) and thereafter, no attempt was made to eliminate any measurements whatsoever. But pits greater than 6 mils in depth were measured wherever measurements could be obtained. When definite pitting occurred but there were no pits greater than 6 mils, the designation P was used. When metal attack occurred on the specimen but no definite pitting could be observed the designation M was used, and when the specimen was unaffected by corrosion, U was used.

Since 1930 it has been the practice to record the 6 deepest pits on all the pipe specimens and the 12 deepest pits on the sheet and plate specimens-6 pits from

each side of the sheet or plate.

The depth gage (fig. 86, A) used at the National Bureau of Standards is an ordinary micrometer depth gage that can be read to the nearest 0.001 inch. The end of the shaft is pointed so as to reach the deepest part of the pit. The base is cut away near the shaft so that the position of the point can be seen, and the originally flat base is machined to a concave cylindrical surface of 4 inch radius, the long axis of the base being parallel to the axis of the cylinder. The gage then will not rock when placed on a 1½-inch or larger pipe surface with the long axis of the base parallel to the pipe axis. The length of the base has also been increased by a bridge for making measurements on severals corredad specimens. by a bridge for making measurements on severely corroded specimens.

The zero setting of the gage must be corrected for the curvature of the pipe surface. This was done by placing the gage on a smooth curved cylinder having the same diameter as the specimen and adjusting the micrometer to read zero. For measuring the pit depths of sheets or plates, the zero reading of the micrometer was adjusted by setting it on a smooth, polished, level surface Ewing [29] has worked out a formula for setting the micrometer on a plane surface and adjusting

the gage for the diameter of the pipe to be measured.

During recent years a different type of micrometer (fig. 86, B) has been used for making the pit-depth measurements, which appears to be less tiresome on the eyes of the observer. The principle of this micrometer is the same as the previously mentioned gage, and the readings can be made to the nearest 0.001 inch on a dial. The base of this gage was machined to a narrow edge, so corrections for the curvature of the pipe were not necessary. A needle extending from the base was moved about in the pit until a maximum reading was obtained on the dial. The zero setting of the gage was adjusted on a smooth, polished, level surface. Check measurements with the two gages gave results that were within the tolerances allowed for the check measurements.

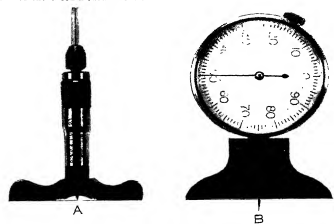


FIGURE 86.—Pit-depth gages.

4. Micrometer gage; B, dial gage. Each reads in thousandth of an inch-mils.

Having accumulated the loss of weight and pit-depth data, the specimens were next labeled and photographed. The pipe specimens were photographed by a circuit camera employing panoramic film (6-inches in width) while the pipes were rotating. In this manner photographs of the specimens 6-inches long were obtained in their actual size. For specimens longer than 6-inches the part showing the severest corrosion was photographed. Pictures were made of all the specimens on which appreciable pitting could be detected. The specimens were then given a coat of colorless varnish to prevent further rusting, and the specimens were preserved at the National Bureau of Standards until 1942, when the demand was made for all available scrap metal for the war effort. All the specimens except the last removals from each set were then turned over to the scrap drive.

#### 2. CALCULATIONS AND METHODS OF REPORTING THE DATA

Usually two specimens of each material were removed from each test site. Hence, the data for losses of weight, rates of loss of weight, maximum penetration, and rates of maximum penetration for the tables involving the National Bureau of Standards tests in section V and to the numerous progress reports referred from the Journal of Research of the National Bureau of Standards are the arithmetical averages of two measurements. When one specimen was missing the value given is the actual value for the loss of weight or maximum penetration of the remaining specimen.

The weighings were always made employing the metric system. Hence, the losses in weight were obtained in grams. For the purpose of comparison, the losses were reduced to ounces per square foot since many of the specimens differed in exposed area. To do this, the loss of weight in grams was multiplied by a factor which gave the loss of weight in ounces per square foot directly. This factor was calculated for each material by the equation

Factor = 
$$\frac{1}{\text{ft}^2 \text{ of exposed area } \times \text{g/oz}} = \frac{1}{28.35A}$$

in which A is the area in square feet. To obtain the rate of loss of weight in ounces per square foot per year the loss in ounces per square foot was divided by the time of burial in years.

The data have often been reported as the average total penetration, which is derived from the loss of weight, the area exposed, and the density of the material. To calculate the average total penetration (in mils) the loss of weight (in ounces per square foot) was multiplied by a factor obtained as follows:

Factor 
$$\frac{12 \times 1000}{\text{wt of ft}^3 \text{ of metal (lb)} \times 16} - \frac{750}{\text{density (lb/ft}^3)}$$

The ratio of the maximum penetration to the average total penetration is the pitting factor. The pitting factor is to some extent a function of the area of pipe surface considered and usually has been found to be somewhat smaller for the 1½-inch specimens than for the 3-inch specimens of similar material.

1½-inch specimens than for the 3-inch specimens of similar material.

The method for obtaining the maximum penetration has already been described. The rate of maximum penetration (in mils per year) is the ratio of the maximum penetration to the time of burial in years. For the purpose of comparing pit depths on pipes of different sizes, the data were reported as the weighted maximum penetration for each material. The exposed area of one 6-inch specimen is equivalent to the exposed area of two 3-inch specimen, and likewise the exposed area of one 3-inch specimen is equivalent to the exposed area of two 1½-inch specimens. Therefore, as there are two specimens for every material, the weighted maximum penetration was obtained as follows: For the 1½-inch specimens the value represents the arithmetical average of the deepest pit on each specimen, a total of two pits for each material. For the 3-inch specimens it represents the arithmetical average of the two maximum pits on each specimen, a total of four pits for each material. For the 6-inch specimens it represents the arithmetical average of the four maximum pits on each specimen, a total of four pits for each of the L and Z materials, as only one specimen of each were taken up at each removal and a total of eight pits for the other 6-inch specimens. The weighted maximum rate of penetration was obtained by dividing the weighted maximum penetration by the time of burial in years.

# XX. APPENDIX 5. CONSTRUCTION OF THE DENISON CORROSION CELL

The dimensions and construction of Denison's most recent modification of his corrosion cell, which differ somewhat from those described in earlier papers, are shown in figure 87. The essential parts of the cell are a bakelite ring 44 mm (1.73 in.) in internal diameter and 21 mm (0.83 in.) high, a metal disk 38 mm (1.50 in.) in diameter, which serves as the anode, another disk of the same material 44 mm (1.73 in.) in diameter, which serves as the cathode and is perforated with 51 holes per square centimeter (329 per square inch), the diameter of each hole being 0.84 mm (0.033 in.). The metal disks are cut with a punch and die from sheet material. A narrow tab, or projection, is provided on the cathode to facilitate electric connection. The total area of the perforated disk is approximately twice the area of one side of the anode.

The other constituents necessary for construction of the cell are the soil samples to be studied which are air-dried and crushed to pass a No. 20 U.S. Standard Sieve, a nonmetallic disk 51 mm (2.00 in.) in diameter containing many perforations and having a hole in the center large enough to permit a calomel electrode to make contact with the adjacent soil, a No. 7 rubber stopper, rubber bands, and a short length of No. 24 copper wire.

The procedure in setting up the cells is as follows. The perforated cathode is cemented in place on a shoulder in the Bakelite ring so that the shoulder becomes a part of the compartment having a height of 10 mm (see fig. 87). This compartment will hereafter be referred to as the shallower compartment or as the lower part of the cell. To fit the cathode in the ring, it is necessary to cut a slit adjacent to the shoulder to allow the projection to pass through. The free space in the slit is sealed with plastic wood. The disk is then cleaned with carbon tetrachloride, using a small stiff fiber brush. The cells in sets of 10 are filled with the soil samples. The shallower compartment is filled first by temporarily placing a disk of filter paper on the reverse side of the cathode to keep the soil from running through the perforations. The cell is then placed on the end of a cylinder about 4 cm high and 4 cm in diameter. The crushed soil is placed in the compartment in increments and packed by placing on the surface of the soil a brass cylinder weighing 500 grams and having a diameter of 44 mm. A disk of filter paper is placed on the soil and moistened with water from a burette. The cell is now inverted, placed on a smooth

⁷The laboratory setup at the National Bureau of Standards was made to accommodate 10 cells. However, a setup may be prepared for a different number of cells.

piece of glass, the filter paper removed, and the lower compartment filled and packed as before. The soil is then saturated by allowing it to absorb water from a strip of filter paper, the cell being supported by 2.5 cm above the surface of the water into which the filter paper dips. The period required for saturation is about 2 hours for most soils. Each cell is placed in an individual dish and covered by a beaker to prevent evaporation. Smoothing of the surface should be avoided after the soil has been moistened.

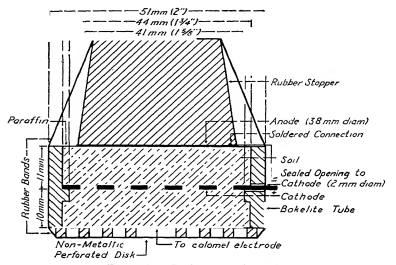


FIGURE 87.—Denison corrosion cell.

The moistening of very heavy soils may be accelerated by adding a slight amount of water to the upper surface of the soil. A sheet of filter paper on the surface of the soil facilitates even distribution of the moisture. The soils are brought under moisture equivalent (the quantity of water retained by a soil under a centrifugal force of 1,000 times gravity) by the method of Schofield [206], as described below. A battery of ten 3-inch (7.6 cm) Buchner funnels set in pressure flasks is prepared to the proper placed in coach funnels as unfaint quantity of this

A battery of ten 3-inch (7.6 cm) Buchner funnels set in pressure flasks is prepared by pouring on a filter paper placed in each funnel a sufficient quantity of a thin suspension of kaolin in water to produce under suction a layer about I mm thick. This layer permits any desired pressure difference up to 1 atmosphere between the interior of the flask and the outer surface of the layer to be maintained. A second piece of filter paper is placed over the kaolin layer and saturated with water, after which the cells are placed in position top down on the paper. A pressure difference of 10 cm of mercury is held for 1 hour, after which it is increased in increments of 5 cm every 10 minutes until a pressure difference of 59 cm is reached. The latter pressure maintained for 1 hour reduces the moisture content of a saturated soil to moisture equivalent. In testing organic soils such as peats, mucks, and tidal marsh, in which the average moisture content in nature is often close to saturation, it is preferable to reduce the moisture content only slightly below saturation rather than to moisture equivalent. This may be accomplished by subjecting the cell to a pressure difference of 5 cm of water for one-half hour.

During the adjustment of the moisture content, the anodes are prepared by rubbing one face with No. IG French emery paper. After cleaning the electrodes with carbon tetrachloride, a No. 24 copper wire is soldered to the reverse side of the anode near the edge.

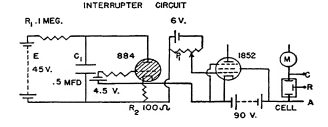
After the soils have reached proper moisture content, the pressure is released and the cells are removed from the funnels. Because shrinkage of the soil results from the withdrawal of water, a crack will often be noticed between the soil and the rim of the Bakelite cylinder. As a crack would provide a channel through which air could pass to the electrodes, it is necessary to seal it with melted paraffin, which is applied by means of a medicine dropper. The anode is placed in position over the

top of the cells and electric connection is made between anode and cathode by joining the copper wire with the projection on the cathode. The perforated insulating disk is placed over the lower end of the cylinder. A No. 7 rubber stopper is then placed on the anode and the whole assembly fastened together by rubber bands. It is especially important that firm contact be maintained between the electrodes and the soil.

The cells are placed in individual 1-pint friction-top cans into which a small amount of water has been poured in order to maintain a saturated atmosphere. In order to prevent a water seal, which would interfere with proper aeration of the cathode when the cell is placed in the can, the cell is inserted in a small glass dish about 4 cm in height and slightly larger than the cell in diameter. The rubber bands around the cell support the cell in the dish in such a way that free circulation of air is insured. A small amount of water is poured into this dish also. After the lids have been sealed, the cans are placed in a thermostatically controlled constant-temperature chamber and maintained at 25° C for 18 to 36 hours in order to stabilize the electrodes. If the room temperature is fairly constant at approximately 25°C, this last step may be omitted. The cells are removed from the cans when the electrical measurements are made, precautions being taken to prevent evaporation of moisture.

# XXI. APPENDIX 6. DARNIELLE'S MODIFICATION OF THE HICKLING APPARATUS FOR MEASURING POTENTIALS

The Hickling method for measuring the potential of polarized electrodes utilizes an electronic interrupter and an electronic potentiometer, by means of which potentials can be measured a very short time after the current has been interrupted.



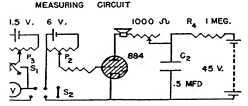


FIGURE 88.—Potentiometer-volt meter and calibration circuit.

Figure 88 shows the essential parts of the interrupter and measuring circuit. Capacitor  $C_1$  is charged through resistor  $R_1$  until the plate of the Thyratron (type 884) is at a certain potential, depending on the grid voltage. The tube then becomes conducting and capacitor  $C_1$  discharges through the tube and resistor  $R_2$  until the capacitor voltage drops to about 15 volts, when the tube becomes nonconducting, and the process is repeated at a rate depending on the values of  $E_1$ ,  $E_1$ ,  $C_2$ , and  $E_2$ .

the process is repeated at a rate depending on the values of  $E_1$ ,  $E_2$ ,  $C_1$ , and  $E_2$ . The cell is in the plate circuit of the pentode (type 1852). Its current may be controlled by potentiometer  $P_1$  and the IE drop across  $E_2$ . Each voltage pulse across  $E_2$  imparts a large negative potential to the grid of the vacuum tube, thereby interrupting the current in the cell. Resistors should be connected in series with the grid of each Thyratron to limit grid current when the tubes conduct current.

The measuring circuit is an oscillating type similar to the interrupter circuit but will not oscillate when the grid of the Thyratron is below a certain critical potential.

In measuring electrode potentials, the electrodes are connected to the grid circuit of a Thyratron, the grid having been adjusted to the critical potential. The cell is so connected that the grid of the Thyratron is most negative between interruptions of the current because of IR drop in the cell. When the current is interrupted, the grid potential will be above or below the critical potential by an amount equal to the

gria potential will be above or below the critical potential by an amount equal to the potential of the electrode. Adjustment of a potentioneter in the grid circuit, so that the critical potential is restored, measures the electrode potential.

In operation, double-pole double-throw switch  $S_2$  is closed to the right (fig. 88), and the grid of the Thyratron in the measuring circuit is adjusted to the critical point by means of potentiometer  $P_2$ , as indicated by slow ticking in the loud-speaker. By reversing switch  $S_2$ , the cell and the voltmeter-potentiometer are introduced into the grid circuit of the Thyratron. If the applied voltage makes the grid more negative than the critical voltage no sound will be heard in the loud-speaker. If the negative than the critical voltage, no sound will be heard in the loud-speaker. If the grid is made more positive than the critical voltage, very rapid ticking will be heard. Potentiometer  $P_3$  is then adjusted until the rate of ticking is identical with the rate when the critical point was adjusted, and the potential difference is read on the voltmeter.

It is important to note that the connections of the measuring circuit to the cell are determined by the directions of IR drop and not by the polarity of the electrodes. Hence in measuring the voltage between the anode and cathode of the cell, the cathode of the cell is connected to the grid. When the voltage between the anode and a reference electrode is measured, the reference electrode is connected to the grid, and for measuring the voltage between the cathode and the reference electrode. the cathode of the cell is connected to the grid. A switching arrangement, not shown in detail in the figure, is used for making rapid changes in connecting to the various electrodes.

When the cell current is interrupted, it drops quickly to a very low value and immediately begins to increase as  $C_1$  discharges. The time required for the current to decrease to a minimum and then increase until the IR drop is more than 0.01 volt. the approximate limit of accuracy of the measurement, may be designated as the "effective" period of interruption. This interval corresponds to the maximum time

during which the electrode depolarizes before its potential is measured.

Hickling's measurements were made with high current densities in cells of very low resistance, a triode being used as the vacuum tube. In order to make the effective period of interruption independent of the polarizing current, the latter was controlled by adjusting the filament current. For measuring potentials in soil cells in which the resistance is high and the current low, the use of a sharp cutoff pentode permitted control of the plate current in the usual way, that is, by regulation of the grid voltage. By eliminating lag in the adjustment of the current in this manner, potential measurements could be made immediately after the current was changed, which is usually desirable in measurements with soil cells. With variation of current and resistance over the extreme range used in corrosion studies, the effective time of interruption was found to be reasonably constant. However, the chief advantage of the pentode over the triode was the greatly improved accuracy of measurement obtained with the former tube. Pentodes of the types 6J7 and 1852 have a very sharp cutoff and provide sufficient plate current for potential measurements in the soil cells. Calculation of the effective period of interruption indicated that this time was about  $8 \times 10^{-5}$  second when  $R_2$  was 75 ohms.

Data show that within the range of current commonly applied to soil-corrosion cells and within the normal range of resistance of these cells, the combined effect of current and resistance on the accuracy of potential measurements is not over 0.01 volt. Measurements accurate to 0.01 volt were made with a current of 4 milliamperes

and a cell resistance of 6,000 ohms.

As the period of interruption of the current is related to the resistance  $R_2$  in the interrupter circuit, varying this resistance provides a convenient means for varying the period of interruption, and hence for studying the rate of depolarization of the soil cells. This was found to be subject to wide variation. In a few soils the change in voltage resulting from a change in  $R_2$  from 100 to 20,000 ohms was little more than 0.01 volt. In other soils this difference was 0.1 volt or more. Rapid depolarization may occur at the anode as well as at the cathode. This period of interruption of the current should be kept as short as possible by keeping  $R_2$  at the lowest practicable value. Measurements of cell potentials by means of the apparatus described have been found to be in good agreement with those by other less generally applicable methods.

# XXII INDEX

Acidity, determination of	
relation to corrosion	2 interpretation
at test sites	2 Depth gages
	2 Depth gages
differential	Differential aereation
corrosion of	39
Aluminum anodes	2 Electrolytic soil resistivity bridge 197
	2 Enamel coatings
corrosion of	Enamel coatings
American Gas Association tests 1	l7
American Petroleum Institute tests 14	
Anodes	1 Field corrosivity tests
aluminum	
carbon 26	8 Galvanized materials, corrosion of 124
magnesium	2 Grease coatings 245
materials 26	32 Grease coatings
noncorrodible 20	25 P
resistance to earth	4 Hickling's apparatus 309
zinc 20	33 Hydrogen-ion concentration, relation to
	1 corrosion 212
control of corrosion	32 relation to total acidity 218
Asbestos cement pipe	28 (
	Ions 11
Bacteria, corrosion caused by174, 19 Bessemer steel	99 + 7 - 3
Bessemer steel	74   Laboratory tests
Bituminous coatings	16 Lead coatings
Bolts, corrosion of	16     Lead coatings     144       15     127       15     varieties tested     34       16     Lite of pipe.     100
Bursting strength of aspestos cement	Tools clamps
effect of corrosion on	97 Leak clamps 106
Cast iron expenses of	Life of pipe
	32
high silicon	o Magnesium Anodes 26
	Malleable cast iron 73
Cathodic areas  control Cathodic protection 2 comparison of methods 2 cont	Materials, composition of 30
control	go corrosion resistant 24:
Cathodic protection 2	50 tested 29
comparison of methods	Mill scale
cost 2	70 Moisture equivalent, definition 23
engineering 2	71
installations 2	01
sources of current 2	
tank bottoms 2	69 Open-hearth iron 7-
2   2   2   2   2   2   2   2   2   2	Oo Orida films on marals
Clay, definition	17 Oxide mins on metals
Cleaning specimens 3	Parkway cable
Clay definition   Cleaning specimens   8	02 Parkway cable
Coating, specifications	48 Pearson coating tester
hituminana 146 9	Pearson-Hadley cathodic protection
concrete 9	46 apparatus
conductivity of	Pipe lines, determination of condition 229
effect of nit denths	68 Pine to soil potential
effect of time	68 Did doubt once relation
inspection	68 Pit depth, area relation
metallic 1	34 time relation
metallic	1
on isolated short sections of pipe 1	59 Prevention of corrosion
reinforced	Protection, criteria for
tests on pipe lines 1	51 Protective current
miscellaneous 1 on isolated short sections of pipe 1 reinforced 152, 156, 1 tests on pipe lines 1 in 1 1	51 Protective current 18 45 Putnam's decomposition potential test 21 98
Columbia rod	98
Conductance test	47 Radio balance resistivity test
Cooperators in corrosion tests	79         Redox apparatus         19           03         Resistivity tests, field         19
Copper, tests of	
copper and copper anoys, analysis of	laboratory
approximate	33 value of
Conner hasring steel	7A Rust, removal of
Conner-molyhdenum iron	74 Scott's pit depth, area, time equation 10
Corfield test	15 Shepard soil resistivity apparatus 19
Copper and copper and spanish of specimens corrosion of 1 Copper bearing steel. Copper-molybdenum iron Corfield test 2 Corrosion, causes estimation of expected 100, 1 literature 2	10 Silt. definition 1
estimation of expected 100. 1	94 Skin effect in cast iron 5
literature	71 Soil acidity
of pipe lines	06 at test sites
potential 1	15   Sinepard son resistivity apparatus   19   Silt, definition   1   194   Skin effect in cast iron   18, 21, 5   18   19   19   19   19   19   19   19
prevention 2	41 definition
products, strength	97 electrical resistivity of
resistant materials 2	42 groups
estimation of expected. 100,1 literature 2 of pipe lines 1 potential 1 prevention 2 products, strength 2 theory 2 time relation 1	85 mechanical analysis of
time relation	90 names
time relation 1 Coupons, pipeline 2 Criteria for protection 2 Current density required for protection 2 Current potential curves 2	85 mechanical analysis of
Current dengity required for protection	51 profiles
Current notantial curves	54 properties vs. corrosivity
Ourrent botential cartes	reports 1
Darnielle's apparatus 3	reports 1 09 resistivity, effect of moisture 21 05 resistivity, effect of pressure 20
Data corresion calculation 3	05 resistivity, effect of pressure 20

# INDEX—Continued

resistivity, effect of temperature 211	Tank bottoms, protection of 269
series 16	Test methods
solution, composition of	Test sites, description of
surveys	Tests of coatings 236
texture	conductivity test 237
tests, comparison of	for pinholes 238
treatment	insulating
types	laboratory
types as indicators of corrosion 199	metallic
Soil corrosivity, determination 194	pattern test
Soil corrosivity tests, Corfield 215	Pearson's holiday detector 23:
coupons 230	reinforcements
correlation, with pit depths 224	Stearn's electronic tester 238
Denison	Theory of corrosion
hydrogen-ion concentration 212	Transmission pipe lines, protection of 261
nipple and can	1
Putnam 214	Volume shrinkage of soils 23
relative merits	1
total acidity	Wenner's four terminal resistivity method. 197
Soils, classification	Wrought iron and steel, composition 30
chemical properties	corrosion of
physical properties	corrosion of
test sites	
Specifications for bituminous coatings 246	Zinc. cleaning SU
Specimens, tested	corrosion of 13
size, effect 56	
Standard error 67	varieties tested 34